



Human ninth edition Genetics

Concepts & Applications

Ricki Lewis



Cell and Molecular Biology

**Human Genetics: Concepts and Applications
9th Edition**

Lewis

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Cell and Molecular Biology

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Preface

Human Genetics for Everyone

Truth is indeed stranger than fiction. When I began writing this textbook 15 years ago with a glimpse of a future where two college roommates take tailored genetic tests, I could never have imagined that today we would be ordering such tests from websites. We send in our DNA on cheek swabs or in saliva samples to learn about our genetic selves. We may receive risk estimates of future health concerns, or take ancestry tests that reveal our pasts, noting which parts of the world our forebears likely came from and maybe even who our distant cousins are. I'm amazed.

Ricki Lewis

Today, human genetics is for everyone. It is about our variation more than about our illnesses, and increasingly about the common rather than the rare. Once an obscure science or an occasional explanation for an odd collection of symptoms, human genetics is now part of everyday conversation. At the same time, it is finally being recognized as the basis of medical science. Despite the popular tendency to talk of “a gene for” this or that, we now know that for most traits and illnesses, several to many genes interact with each other and environmental influences. By coming to know our genetic backgrounds, we can control our environments in more healthful ways. Genetic knowledge is, therefore, both informative and empowering. This book shows you how and why this is true.

What Sets this Book Apart

Current Content

As a member of the Information and Education Committee of the American Society of Human Genetics, an instructor of “Genethics,” genetic counselor, and long-time science writer, Dr. Lewis is aware of research news and government policy changes before they are published. The most exciting new developments find their way into each edition of *Human Genetics: Concepts and Applications*, sometimes in the words of the people they directly affect. A few of the most compelling updates to this edition include

- Direct-to-consumer genetic testing
- Genome-wide association (GWA) studies: promises and perils
- Gene expression profiling and personalized medicine
- Human microbiome project
- Human variation and ancestry
- GINA (Genetic Information Nondiscrimination Act)
- Induced pluripotent stem cells (reprogramming)

This new edition also reflects the shift in focus in the field of human genetics from rare single-gene inheritance to more common multifactorial traits and disorders.

The Human Touch

Human genetics is about people, and their voices echo throughout these pages. Most are real, some are composites, and many are based on the author’s experience as a science writer, genetic counselor, and hospice volunteer.

Compelling Stories and Case Studies Lewis enlivens her clear presentation of genetic concepts with compelling stories and cases like the following:

- A young fashion magazine editor keeping her leukemia at bay thanks to a drug developed through genetic research (Ch. 18, p. 366)
- A man freed from a 25-year prison term following reconsideration of DNA evidence (Ch. 14, p. 265)
- A father whose little girl has a condition so rare that it doesn’t even have a name (Ch. 4, p. 69)

Practical Application of Human Genetics Recognizing that the goal of most introductory science courses is to better inform future voters and consumers, the author provides practical application of the content to students’ lives. Topics of particular interest to students include:

- The role that genes play in disease susceptibility, physical characteristics, body weight, and behaviors, with an eye toward the dangers of genetic determinism
- Biotechnologies, including genetic testing, gene therapy, stem cell therapy, gene expression profiling, genome-wide association studies, and personalized medicine
- Ethical concerns that arise from the interface of genetic information and privacy, such as infidelity testing, ancestry testing, and direct-to-consumer genetic testing

The Lewis Guided Learning System

Each chapter is framed with a set of pedagogical features designed to reinforce the key ideas in the chapter and prompt students to think more deeply about the application of the content they have just read.

Dynamic Art

Outstanding photographs and dimensional illustrations, vibrantly colored, are featured throughout *Human Genetics*. Students will learn from a variety of figure types, including process figures with numbered steps, micro to macro representations, and the combination of art and photos to relate stylized drawings to real-life structures.

New to this Edition!

New and updated information is integrated throughout the chapters, and a few features from past editions have been moved. Highlights from the revision are included here.

Chapter 1 *Overview of Genetics*

- Updates on the Genetic Information Nondiscrimination Act and the Human Microbiome Project
- New Figure 1.8 Diseaseome—diseases are connected in unexpected ways
- New *Bioethics: Choices for the Future*, “Genetic Testing and Privacy”

Chapter 2 *Cells*

- Stem cell coverage now stresses reprogrammed cells, with two new figures and a new *Bioethics: Choices for the Future*, “Should You Bank Your Stem Cells?”
- New *In Their Own Words*, “A Little Girl with Giant Axons”

Chapter 4 *Single-gene Inheritance*

- New chapter opener “His Daughter’s DNA,” about a father’s quest to solve a genetic mystery
- New section 4.1, A Tale of Two Families

Chapter 5 *Beyond Mendel’s Laws*

- New Reading 5.1, “The Genetic Roots of Alzheimer Disease”
- New Table 5.3, Types of Genetic Markers

Chapter 6 *Matters of Sex*

- New chapter opener, “A Controversial Hypothesis: Mental Illness, Mom, and Dad”
- New Reading 6.2, “Rett Syndrome—A Curious Inheritance Pattern”

Chapter 7 *Multifactorial Traits*

- New Figure 7.1, Anatomy of a trait—rare single-gene disorders versus common SNP patterns
- New section 7.4, Genome-wide association studies (including new figures 7.9 and 7.11)

Chapter 8 *Genetics of Behavior*

- New section 8.5, How nicotine is addictive and raises cancer risk
- New section 8.8, Autism (includes new Figure 8.9, Understanding autism)

Chapter 9 *DNA Structure and Replication*

- New *Bioethics: Choices for the Future*, “Infidelity Testing”

Chapter 11 *Gene Expression and Epigenetics*

- New Figure 11.7, Control of gene expression (transcription factors and microRNAs)
- New text on the evolving definition of a gene

Chapter 12 *Gene Mutation*

- New chapter opening case study, “The Amerithrax Story”
- New Figure 12.1, Animal models of human diseases
- New Figure 12.11, Using copy number variants in healthcare

Chapter 13 *Chromosomes*

- New *Bioethics: Choices for the Future*, “The Denmark Study: Screening for Down Syndrome”

Chapter 16 *Human Ancestry*

- New *Bioethics: Choices for the Future*, “Indigenous Peoples”
- Expanded coverage of markers, haplogroups, and migration
- New Reading 16.2 “Should You Take a Genetic Ancestry Test?”

Chapter 17 *Genetics of Immunity*

- Shortened and reorganized to stress genetics

Chapter 18 *Genetics of Cancer*

- New Table 18.2, Processes and Pathways Affected in Cancer
- The cancer genome

Chapter 19 *Genetic Technologies: Amplifying, Modifying, and Monitoring DNA*

- Expanded and updated information on DNA patents
- New section 19.5, Silencing DNA (RNAi, antisense, and knockouts)

Chapter 20 *Genetic Testing and Treatment*

- New section 20.1, “Geneticists find zebras, and some horses” (including new figure 20.1)
- New information on direct-to-consumer tests and CLIA regulations
- Gene therapy to treat hereditary blindness in an 8-year-old

Chapter 22 *Genomics*

- New chapter opener, “20,000 Genomes and Counting”
- New Reading 22.1, “The First Three Humans to Have Their Genomes Sequenced”

PART 1 Introduction



Personal genetic information is now readily available. People use genetic information to learn about their health risks and trace their ancestry.

CHAPTER

1

Overview of Genetics

Direct-to-Consumer Genetic Testing

Genetic tests were once used solely to diagnose conditions so rare that doctors could not often match a patient's symptoms to a recognized illness. Today, taking a genetic test is as simple as ordering a kit on the Internet, swishing a plastic swab inside the mouth, and mailing the collected cell sample to a testing company or research project. The returned information can reach back to the past to chart a person's ancestry, or into the future to estimate disease risk.

Some "direct-to-consumer" (dTC) genetic tests identify well-studied mutations that cause certain diseases. Yet other tests are based on "associations" of patterns of genetic variation that appear in people who share certain traits or illnesses, but not nearly as often in others. Because these new types of tests are drawn from population studies, they might not apply to a particular person. Consumers who take Internet-offered tests can review results with a genetic counselor. If interpreted carefully, information from genetic tests can be used to promote health or identify relatives.

Eve is curious about her ancestry and future health, so she finds a company whose tests provide clues to both. Her DNA sample is scanned for variants inherited from her mother against a database of patterns from 20 nations and 200 ethnic groups in and near Africa. Eve learns that her family on her mother's side came from Gambia. She will be notified of others who share this part of her deep ancestral roots.

Chapter Contents

1.1 Introducing Genes

1.2 Levels of Genetics

The Instructions: DNA, Genes, Chromosomes, and Genomes

The Body: Cells, Tissues, and Organs

Relationships: From Individuals to Families

The Bigger Picture: From Populations to Evolution

1.3 Genes and Their Environment

1.4 Applications of Genetics

Establishing Identity

Health Care

Agriculture

Ecology

A Global Perspective

The health tests require more thought. Eve dismisses tests for traits she considers frivolous—ear wax consistency and ability to taste bitter foods—as well as for the obvious, such as blue eyes, baldness, or obesity. She already knows if she overeats and doesn't exercise, she'll gain weight. Cancer and Alzheimer disease are too remote for a 20-year-old to think much about, so she foregoes those tests too—for now.

Eve selects her health tests based on her family history—she, a sister, and her father often have respiratory infections. So she asks for her DNA to be tested for gene variants that might affect breathing—cystic fibrosis, asthma, emphysema, nicotine dependence, and lung cancer. Reluctantly she checks the boxes for heart and blood vessel diseases, too. Her reasoning: She can do something proactive to prevent or delay these conditions, such as breathing clean air, exercising, not smoking, and following a healthy diet.

Is genetic testing something that you would do?

1.1 Introducing Genes

Genetics is the study of inherited traits and their variation. Sometimes people confuse genetics with genealogy, which considers relationships but not traits. With the advent of tests that can predict genetic illness, genetics has even been compared to fortunetelling! But genetics is neither genealogy nor fortunetelling—it is a life science.

Inherited traits range from obvious physical characteristics, such as the freckles and red hair of the girl in **figure 1.1**, to many aspects of health, including disease. Talents, quirks, behaviors, and other difficult-to-define characteristics might appear to be inherited if they affect several family members, but may reflect a combination of genetic and environmental influences. Some traits attributed to genetics border on the silly—such as sense of humor, fondness for sports, and whether or not one votes.

Until the 1990s, genetics was more an academic than a clinical science, except for rare diseases inherited in clear patterns in families. As the century drew to a close, researchers completed the global Human Genome Project, which deciphered the complete set of our genetic instructions. The next step—surveying our genetic variability—was already underway. Today, genetics has emerged as an informational as well as a life science that is having a huge societal impact. Genetic information is accessible to anyone, and the contribution of genes to the most common traits and disorders is increasingly appreciated.

Like all sciences, genetics has its own vocabulary. Many terms may be familiar, but actually have precise technical definitions. All of the terms and concepts in this chapter are merely introductions that set the stage for the detail in subsequent chapters.

Genes are the units of heredity, which is the transmission of inherited traits. Genes are biochemical instructions that tell



Figure 1.1 Inherited traits. This young lady owes her red hair, fair skin, and freckles to a variant of a gene that encodes a protein (the melanocortin 1 receptor) that controls the balance of pigments in the skin.

cells, the basic units of life, how to manufacture certain proteins. These proteins, in turn, impart or control the characteristics that create much of our individuality. A gene is the long molecule **deoxyribonucleic acid (DNA)**. It is the DNA that transmits information, in its sequence of four types of building blocks.

The complete set of genetic instructions characteristic of an organism, including protein-encoding genes and other DNA sequences, constitutes a **genome**. Nearly all of our cells contain two copies of the genome. Researchers are still analyzing what all of our genes do, and how genes interact and respond to environmental stimuli. Only a tiny fraction of the 3.2 billion building blocks of our genetic instructions determines the most interesting parts of ourselves—our differences. Comparing and analyzing genomes, which constitute the field of **genomics**, reveals how closely related we are to each other and to other species.

Genetics directly affects our lives, as well as those of our relatives, including our descendants. Principles of genetics also touch history, politics, economics, sociology, art, and psychology. Genetic questions force us to wrestle with concepts of benefit and risk, even tapping our deepest feelings about right and wrong. A field of study called **bioethics** was founded in the 1970s to address moral issues and controversies that arise in applying medical technology. Bioethicists today confront concerns that new genetic knowledge raises, such as privacy and discrimination. Essays throughout this book address bioethical issues.

Many of the basic principles of genetics were discovered before DNA was recognized as the genetic material, from experiments and observations on patterns of trait transmission in families. For many years, genetics textbooks (such as this one) presented concepts in the order that they were understood, discussing pea plant experiments before DNA structure. Now, since even gradeschoolers know what DNA is, a “sneak preview” of DNA structure and function is appropriate (**Reading 1.1**) to consider the early discoveries in genetics (chapter 4) from a modern perspective.

Reading 1.1

Introducing DNA

We have probably wondered about heredity since our beginnings, when our distant ancestors noticed family traits such as a beaked nose or an unusual skill, such as running fast or manual dexterity. Awareness of heredity appears in ancient Jewish law that excuses a boy from circumcision if his brothers or cousins bled to death following the ritual. Nineteenth-century biologists thought that body parts controlled traits, and they gave the hypothetical units of inheritance such colorful names as “pangens,” “ideoblasts,” “gemules,” and simply “characters.”

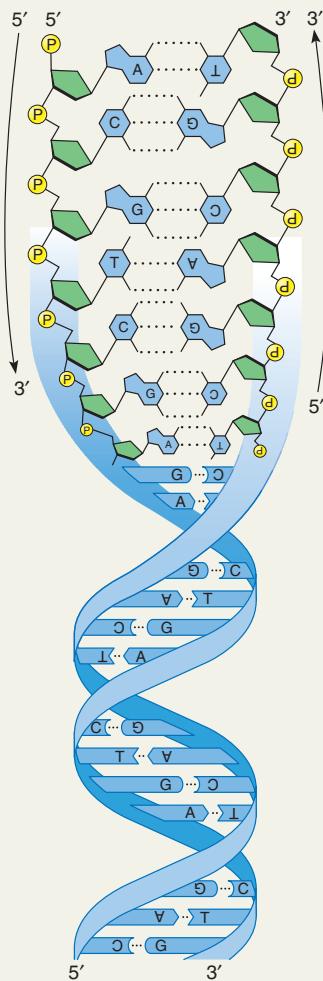


Figure 1 The DNA double helix. The 5' and 3' labels indicate the head-to-tail organization of the DNA double helix. A, C, T, and G are bases. S stands for sugar and P for phosphate.

In the late nineteenth century, when Gregor Mendel bred pea plants to follow trait transmission, establishing the basic laws of inheritance, he inferred that units of inheritance were at play. He had no knowledge of cells, chromosomes, or DNA. This short reading explains, very briefly, what Mendel did not know—how DNA confers inherited traits. Chapter 9 examines DNA in detail.

DNA resembles a spiral staircase or double helix in which the “rails” or backbone of alternating sugars and phosphates is the same from molecule to molecule, but the “steps” are pairs of four types of building blocks, or DNA bases, whose sequence varies (**figure 1**). The chemical groups that form the steps are adenine (A) and thymine (T), which attract, and cytosine (C) and guanine (G), which attract. DNA holds information in the sequences of A, T, C, and G. The two strands are oriented in opposite directions.

DNA uses its information in two ways. If the sides of the helix part, each half can reassemble its other side by pulling in free building blocks—A and T attracting and G and C attracting. This process, called DNA replication, maintains the information when the cell divides. DNA also directs the production of specific proteins. In a process called transcription, the sequence of part of one strand of a DNA molecule is copied into a related molecule, messenger RNA. Each three such RNA bases in a row attract another type of RNA that functions as a connector, bringing with it a particular amino acid, which is a building block of protein. The synthesis of a protein is called translation. As the two types of RNA temporarily bond, the amino acids align and join, forming a protein that is then released. DNA, RNA, and proteins can be thought of as three related languages of life (**figure 2**).

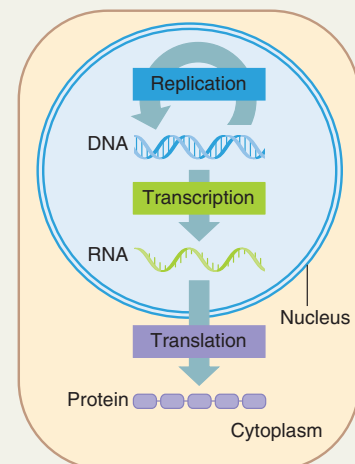


Figure 2 The language of life: DNA to RNA to protein.

1.2 Levels of Genetics

Genetics considers the transmission of information at several levels. It begins with the molecular level and broadens through cells, tissues and organs, individuals, families, and finally to populations and the evolution of species (**figure 1.2**).

The Instructions: DNA, Genes, Chromosomes, and Genomes

Genes consist of sequences of four types of DNA building blocks, or bases—adenine, guanine, cytosine, and thymine, abbreviated A, G, C, and T. Each base bonds to a sugar and a phosphate group to form a unit called a nucleotide, and nucleotides are linked into long DNA molecules. In genes, DNA bases provide an alphabet of sorts. Each consecutive three DNA bases is a code for a particular amino acid, and amino acids are the building blocks of proteins. Another type of molecule, **ribonucleic acid (RNA)**, uses the information in certain DNA sequences to construct specific proteins. Messenger RNA (mRNA) carries the gene's base sequence, whereas two other major types of RNA assemble the protein's building blocks. These proteins confer the trait. DNA remains in the part of the cell called the nucleus, and is passed on when a cell divides.

Proteomics is a field that considers the types of proteins made in a particular type of cell. A muscle cell, for example, requires abundant contractile proteins, whereas a skin cell contains mostly scaly proteins called keratins. A cell's proteomic profile changes as conditions change. A cell lining the stomach, for example, would produce more protein-based digestive enzymes after a meal.

The human genome has about 20,325 protein-encoding genes. The few thousand known to cause disorders or traits are

described in a database called Online Mendelian Inheritance in Man (MIM). It can be accessed through the National Center for Biotechnology Information (<http://www.ncbi.nlm.nih.gov/>). Throughout this text, the first mention of a disease includes its MIM number. **Reading 4.1** describes some of the more colorful traits in MIM.

Despite knowing the sequence of DNA bases of the human genome, there is much we still do not know. For example, only about 1.5 percent of our DNA encodes protein. The rest includes many highly repeated sequences that assist in protein synthesis or turn protein-encoding genes on or off, and other sequences whose roles are yet to be discovered.

The same protein-encoding gene may vary slightly in base sequence from person to person. These variants of a gene are called **alleles**. The changes in DNA sequence that distinguish alleles arise by a process called **mutation**. Once a gene mutates, the change is passed on when the cell that contains it divides. If the change is in a sperm or egg cell that becomes a fertilized egg, it is passed to the next generation.

Some mutations cause disease, and others provide variation, such as freckled skin. Mutations can also help. For example, a mutation makes a person's cells unable to manufacture a surface protein that binds HIV. These people are resistant to HIV infection. Many mutations have no visible effect because they do not change the encoded protein in a way that affects its function, just as a minor spelling *error* does not obscure the meaning of a sentence.

Parts of the DNA sequence can vary among individuals, yet not change appearance or health. Such a variant in sequence that is present in at least 1 percent of a population is called a **polymorphism**, which means "many forms." The genome includes millions of single base sites that differ among individuals. These are called **single nucleotide polymorphisms**

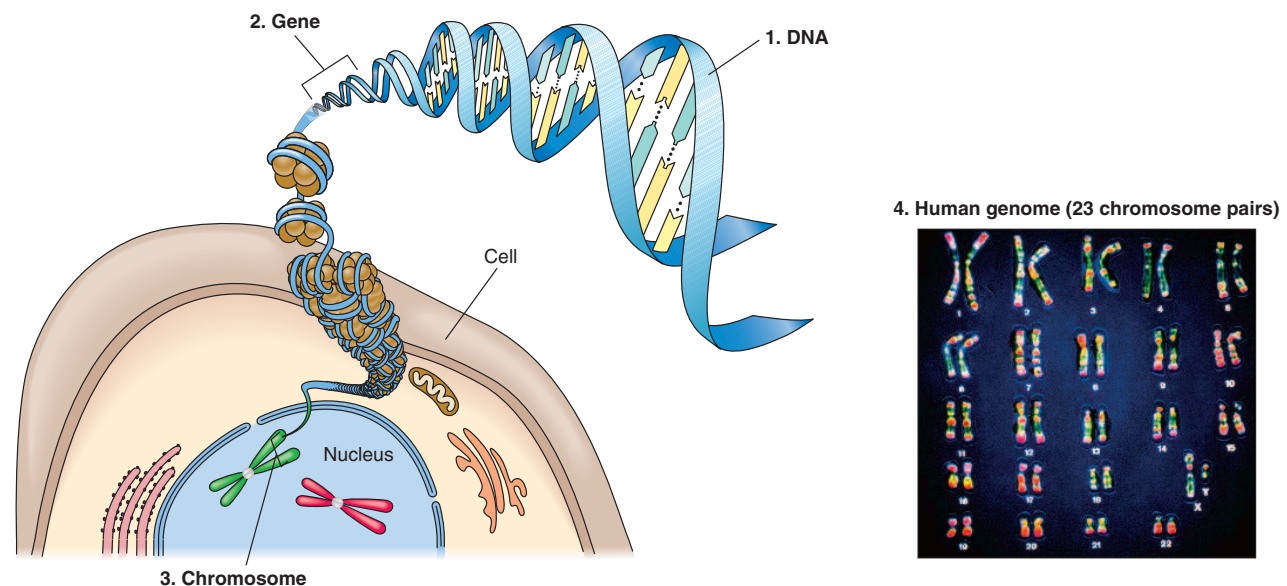


Figure 1.2 Levels of genetics. Genetics can be considered at several levels, from DNA, to genes, to chromosomes, to genomes, to the more familiar individuals, families, and populations. (A gene is actually several hundred or thousand DNA bases long.)

(SNPs, pronounced “snips”). SNPs can cause disease or just mark places in the genome where people differ.

Many research groups are conducting **genome-wide association studies** that look at SNPs in thousands of individuals to identify and track combinations of these landmarks of genetic variation that are found almost exclusively among people with a particular disorder or trait. These SNP patterns can then be used to estimate risk of the disease in people who are not yet sick but have inherited the same DNA variants.

The information in the human genome is studied in several ways, and at several levels. The DNA base sequence can be deciphered for a specific gene that causes a specific illness. Deducing the encoded protein’s structure and function by searching gene-protein databases for similar sequences may explain the symptoms. At the other end of the informational spectrum is sequencing an entire genome. A genome-wide association study lies in between the sequencing of a gene and a genome in scope. If a genome is like a detailed Google map of the entire United States and a gene is like a Google map showing the streets of a neighborhood, then SNPs that speckle a genome are like a map of the United States with only the names of states and interstate highways indicated—just clues.

Sequences of DNA bases, whether for single genes or entire genomes, provide a structural view of genetic material. Another way to look at DNA, called **gene expression profiling**, highlights function by measuring the abundance of different RNA molecules in a cell. These RNAs reflect protein production. In this way, gene expression profiles showcase a cell’s activities. The power of the approach is in comparisons. A muscle cell from a bedridden person, for example, would have different levels of contractile proteins than the same type of cell from an active athlete. **Table 1.1** summarizes types of information that DNA sequences provide.

DNA molecules are very long. They wrap around proteins and wind tightly, forming structures called **chromosomes**. A human somatic (non-sex) cell has 23 pairs of chromosomes. Twenty-two pairs are **autosomes**, which do not differ between the sexes. The autosomes are numbered from 1 to 22, with 1 the largest. The other two chromosomes, the X and the Y, are **sex chromosomes**. The Y chromosome bears genes that determine

maleness. In humans, a female has two X chromosomes and a male has one X and one Y. Charts called **karyotypes** display the chromosome pairs from largest to smallest.

A human cell has two complete sets of genetic information. The 20,325 or more protein-encoding genes are scattered among 3.2 billion DNA bases in each set of 23 chromosomes.

The Body: Cells, Tissues, and Organs

A human body consists of approximately 50 to 100 trillion cells. All cells except red blood cells contain the entire genome, but cells differ in appearance and activities because they use only some of their genes—and which ones they access at any given time depends upon environmental conditions both inside and outside the body.

The genome is like the Internet in that it contains a wealth of information, but only some of it need be accessed. The expression of different subsets of genes drives the **differentiation**, or specialization, of distinctive cell types. An adipose cell is filled with fat, but not the scaly keratins that fill skin cells, or the collagen and elastin proteins of connective tissue cells. All three of these cell types, however, have complete genomes. Groups of differentiated cells assemble and interact with each other and the nonliving material that they secrete to form aggregates called tissues.

The body has only four basic tissue types, composed of more than 260 types of cells. Tissues intertwine and layer to form the organs of the body, which in turn connect into organ systems. The stomach shown at the center of **figure 1.3**, for example, is a sac made of muscle that also has a lining of epithelial tissue, nervous tissue, and a supply of blood, which is a type of connective tissue. **Table 1.2** describes tissue types.

Many organs include rare, unspecialized **stem cells**. A stem cell can divide to yield another stem cell and a cell that differentiates. Thanks to stem cells, organs can maintain a reserve supply of cells to grow and repair damage.

Relationships: From Individuals to Families

Two terms distinguish the alleles that are *present* in an individual from the alleles that are *expressed*. The **genotype** refers to the underlying instructions (alleles present), whereas the **phenotype** is the visible trait, biochemical change, or effect on health (alleles expressed). Alleles are further distinguished by how many copies it takes to affect the phenotype. A **dominant** allele has an effect when present in just one copy (on one chromosome), whereas a **recessive** allele must be present on both chromosomes to be expressed.

Individuals are genetically connected into families. A person has half of his or her genes in common with each parent and each sibling, and one-quarter with each grandparent. First cousins share one-eighth of their genes.

For many years, transmission (or Mendelian) genetics dealt with single genes in families. The scope of transmission genetics has greatly broadened in recent years. Family genetic studies today often trace more than one gene at a time, or traits that have substantial environmental components. Molecular genetics, which considers DNA, RNA, and proteins, often

Table 1.1 Types of Information in DNA Sequences

Level	Description
Single gene	Hundreds to thousands of DNA bases that encode a protein or parts of a protein
Genome	The entire 3.2-billion base sequence of the genetic material in a human cell
Genome-wide association study	Patterns of single-base variants (SNPs) correlated to traits or medical conditions
Gene expression profiling	Levels of mRNAs in specific cells under specific conditions that reflect physiology and reveal abnormalities in function

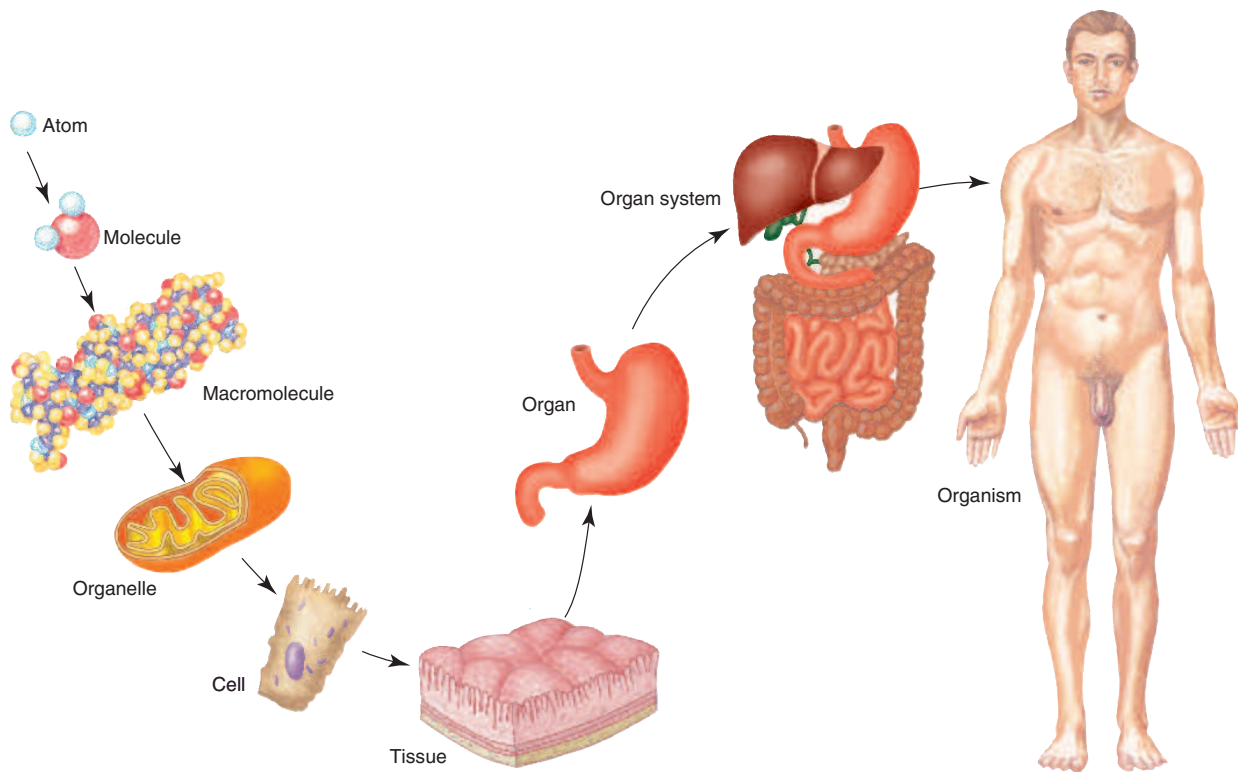


Figure 1.3 Levels of biological organization.

Table 1.2 Tissue Types	
Tissue	Function/Location/Description
Connective tissues	A variety of cell types and materials around them that protect, support, bind to cells and fill spaces throughout the body; include cartilage, bone, blood, and fat
Epithelium	Tight cell layers that form linings that protect, secrete, absorb, and excrete
Muscle	Cells that contract, providing movement
Nervous	Neurons transmit information as electrochemical impulses that coordinate movement and sense and respond to environmental stimuli; neuroglia are cells that support and nourish neurons

begins with transmission genetics, when an interesting family trait or illness comes to a researcher's attention. Charts called **pedigrees** represent the members of a family and indicate which individuals have particular inherited traits. Chapter 4 includes many pedigrees.

Sometimes understanding a rare condition inherited as a single-gene trait leads to treatments for the greater number of people with similar disorders that are not inherited. This is the case for the statin drugs widely used to lower cholesterol. Still, despite the availability of the human genome sequence, some single-gene

disorders are so rare that they do not even have a name. The opening essay to chapter 4 describes a little girl in this situation.

The Bigger Picture: From Populations to Evolution

Above the family level of genetic organization is the population. In a strict biological sense, a population is a group of interbreeding individuals. In a genetic sense, a population is a large collection of alleles, distinguished by their frequencies. People from a Swedish population, for example, would have a greater frequency of alleles that specify light hair and skin than people from a population in Ethiopia, who tend to have dark hair and skin. The fact that groups of people look different and may suffer from different health problems reflects the frequencies of their distinctive sets of alleles. All the alleles in a population constitute the **gene pool**. (An individual does not have a gene pool.)

Population genetics is applied in health care, forensics, and other fields. It is also the basis of evolution, which is defined as changing allele frequencies in populations. These small-scale genetic changes foster the more obvious species distinctions we most often associate with evolution.

Comparing DNA sequences for individual genes, or the amino acid sequences of the proteins that the genes encode, can reveal how closely related different types of organisms are (**figure 1.4**). The underlying assumption is that the more similar the sequences are, the more recently two species diverged from a shared ancestor. This is a more plausible explanation than two species having evolved similar or identical gene sequences by chance.

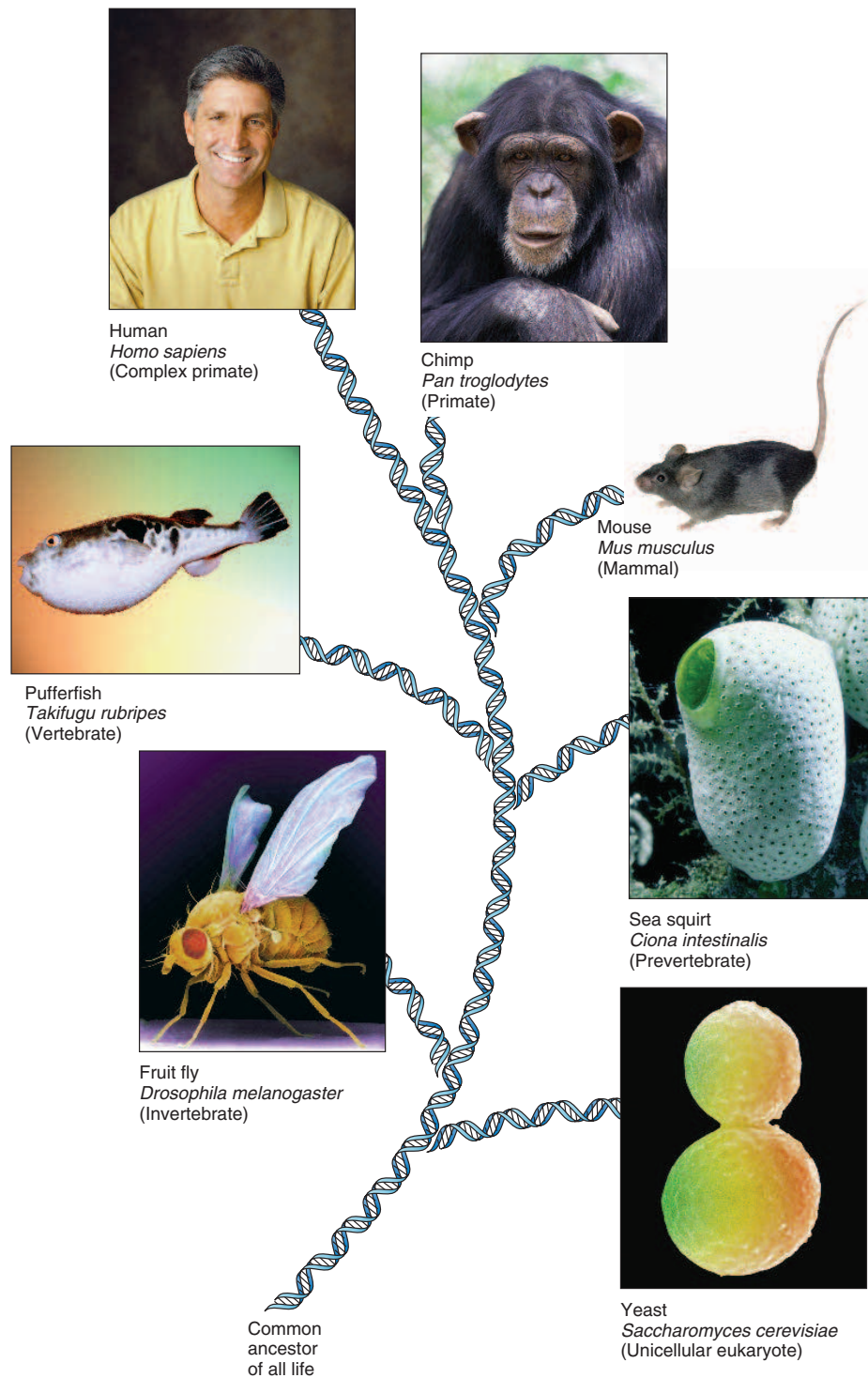


Figure 1.4 Genes and genomes reveal our place in the world. All life is related, and different species share a basic set of genes that makes life possible. The more closely related we are to another species, the more genes we have in common. This illustration depicts how humans are related to certain contemporaries whose genomes have been sequenced.

During evolution, species diverged from shared ancestors. For example, humans diverged more recently from chimps, our closest relative, than from mice, pufferfish, sea squirts, flies, or yeast.

Both the evolution of species and family patterns of inherited traits show divergence from shared ancestors. This is based on logic. It is more likely that a brother and sister share approximately half of their gene variants because they have the same parents than that half of their genetic material is identical by chance.

Genome sequence comparisons reveal more about evolutionary relationships than comparing single genes, simply because there are more data. Humans, for example, share more than 98 percent of the DNA sequence with chimpanzees. Our genomes differ from theirs more in gene organization and in the number of copies of genes than in the overall sequence. Learning the functions of the human-specific genes may explain the differences between us and them—such as our lack of hair and use of spoken language. Reading 16.1 highlights some of our distinctively human traits.

At the level of genetic instructions for building a body, we are not very different from other organisms. Humans also share many DNA sequences with mice, pufferfish, and fruit flies. Dogs get many of the same genetic diseases that we do! We even share some genes necessary for life with simple organisms such as yeast and bacteria.

Comparisons of people at the genome level reveal that we are much more like each other genetically than are other mammals. It's odd to think that chimpanzees are more distinct from each other than we are! The most genetically diverse modern people are from Africa, where humanity arose. The gene variants among different modern ethnic groups include subsets of our ancestral African gene pool.

Key Concepts

1. Genetics is the study of inherited traits and their variation.
2. Genetics can be considered at the levels of DNA, genes, chromosomes, genomes, cells, tissues, organs, individuals, families, and populations.
3. A gene can exist in more than one form, or allele.
4. Comparing genomes among species reveals evolutionary relatedness.

1.3 Genes and Their Environment

Despite the focus of genetics on single-gene traits for many years, nearly all genes do not function alone but are influenced by the actions of other genes, as well as by factors in the environment. For example, a number of genes control how much energy (calories) we extract from food. However, the numbers and types of bacteria that live in our intestines vary from person to person, and affect how many calories we extract from food. This is one reason why some people can eat a great deal and not gain weight, yet others gain weight easily. Studies show that an item of food—such as a 110-calorie cookie—may yield 110 calories in one person's body, but only 90 in another's.

Multifactorial, or complex, traits are those that are determined by one or more genes and the environment (**figure 1.5**).

(The term *complex traits* has different meanings in a scientific and a popular sense, so this book uses the more precise term *multifactorial*.) The same symptoms may be inherited or not, and if inherited, may be caused by one gene or more than one. Usually the inherited forms of an illness are rarer, as is the case for Alzheimer disease, breast cancer, and Parkinson disease.

Knowing whether a trait or illness is single-gene or multifactorial is important for predicting the risk of occurrence in a particular family member. This is simple to calculate using the laws that Mendel derived, discussed in chapter 4. In contrast, predicting the recurrence of a multifactorial trait or disorder in a family is difficult because several contributing factors are at play.

Osteoporosis illustrates the various factors that can contribute to a disease. It mostly affects women past menopause, thinning the bones and increasing risk of fractures. Several genes contribute to susceptibility to the condition, as well as do lifestyle factors, including smoking, lack of weight-bearing exercise, and a calcium-poor diet.

The modifying effect of the environment on gene action counters the idea of **genetic determinism**, which is that an inherited trait is inevitable. The idea that “we are our genes,” or such phrases as “its in her DNA,” dismiss environmental influences. In predictive testing for inherited disease, which detects a disease-causing genotype in a person without symptoms, results are presented as risks, rather than foregone conclusions, because the environment can modify gene expression. A woman might be told “You have a 45 percent chance of developing this form of breast cancer,” not, “You will get breast cancer.”

Genetic determinism may be harmful or helpful, depending upon how we apply it. As part of social policy, genetic determinism can be disastrous. An assumption that one ethnic group is genetically less intelligent than another can lead to lowered expectations and/or fewer educational opportunities for those perceived as biologically inferior. Environment, in fact, has a huge impact on intellectual development.

Identifying the genetic component to a trait can, however, be helpful in that it gives us more control over our health by guiding us in influencing noninherited factors, such as diet. This is the case for the gene that encodes a liver enzyme called hepatic lipase. It controls the effects of eating a fatty diet by regulating the balance of LDL (“bad cholesterol”) to HDL (“good cholesterol”) in the blood after such a meal. Inherit one allele and a person can eat a fatty diet yet have a healthy cholesterol profile. Inherit a different allele and a slice of chocolate cake or a fatty burger sends LDL up and HDL down—an unhealthy cholesterol profile.

Key Concepts

1. Inherited traits are determined by one gene (Mendelian) or by one or more genes and the environment (multifactorial).
2. Even the expression of single genes is affected to some extent by the actions of other genes.
3. Genetic determinism is the idea that an inherited trait cannot be modified.



a.



b.

Figure 1.5 Mendelian versus multifactorial traits. (a) Polydactyly—extra fingers and/or toes—is a Mendelian trait (single-gene). (b) Hair color is multifactorial, controlled by at least three genes plus environmental factors such as the bleaching effects of sun exposure.

1.4 Applications of Genetics

Barely a day goes by without some mention of genetics in the news. Genetics is impacting many areas of our lives, from health care choices, to what we eat and wear, to unraveling our pasts and controlling our futures. Thinking about genetics evokes fear, hope, anger, and wonder, depending on context and circumstance. Following are glimpses of applications of genetics that we will explore more fully in subsequent chapters.

Establishing Identity

Comparing DNA sequences to establish or rule out identity, relationships, or ancestry is becoming routine. This approach, called DNA profiling, looks at SNPs and short, repeated DNA sequences. It has many applications.

Forensics

Before September 11, 2001, the media reported on DNA profiling (then known as DNA fingerprinting) rarely, usually to identify plane crash victims or to provide evidence in high-profile criminal cases. After the 2001 terrorist attacks, investigators compared DNA sequences in bones and teeth collected from the scenes to hair and skin samples from hairbrushes, toothbrushes, and clothing of missing people, and to DNA samples from relatives. It was a massive undertaking that would soon be eclipsed by natural disasters such as the need to identify victims of the tsunami in Asia in 2004 and hurricane Katrina in the United States in 2005.

A more conventional forensic application matches a rare DNA sequence in tissue left at a crime scene to that of a sample from a suspect. This is statistically strong evidence that the accused person was at the crime scene (or that someone planted evidence). DNA databases of convicted felons often provide “cold hits” when DNA at a crime scene matches a criminal’s DNA in the database. This is especially helpful when there is no suspect.

DNA profiling is used to overturn convictions, too. Illinois led the way in 1996, when DNA tests exonerated the Ford Heights Four—men convicted of a gang rape and double murder who had spent 18 years in prison, 2 of them on death row. In 1999,

the men received compensation of \$36 million for their wrongful convictions. A journalism class at Northwestern University initiated the investigation that gained the men their freedom. The case led to new state laws granting death row inmates new DNA tests if their convictions could have arisen from mistaken identity, or if DNA tests were performed when they were far less accurate. The Innocence Project is an organization that has used DNA profiling to exonerate more than 200 death row prisoners. One of them is introduced in the opening essay to chapter 14.

DNA profiling helps adopted individuals locate blood relatives. The Kinsearch Registry maintains a database of DNA information on people adopted in the United States from China, Russia, Guatemala, and South Korea, which are the sources of most foreign adoptions. Adopted individuals can provide a DNA sample and search the database by country of origin to find siblings. Websites allow children of sperm donors to find their biological fathers, if the men wish to be contacted.

History and Ancestry

DNA analysis can help to flesh out details of history. Consider the offspring of Thomas Jefferson’s slave, Sally Hemings (figure 1.6). Rumor at the time placed Jefferson near Hemings nine months before each of her seven children was born, and the children themselves claimed to be presidential offspring. A Y chromosome analysis revealed that Thomas Jefferson could have fathered Hemings’s youngest son, Eston—but so could any of 26 other Jefferson family members. The Y chromosome, because it is only in males, passes from father to son. Researchers identified very unusual DNA sequences on the Y chromosomes of descendants of Thomas Jefferson’s paternal uncle, Field Jefferson. (These men were checked because the president’s only son with wife Martha died in infancy, so he had no direct descendants.) The Jefferson family’s unusual Y chromosome matched that of descendants of Eston Hemings, supporting the talk of the time.

Reaching farther back, DNA profiling can clarify relationships from Biblical times. Consider a small group of Jewish people, the cohanim, who share distinctive Y chromosome DNA sequences and enjoy special status as priests. By considering the number of DNA differences between cohanim and other



Figure 1.6 DNA reveals and clarifies history. After DNA evidence showed that Thomas Jefferson likely fathered a son of his slave, descendants of both sides of the family met.

Jewish people, how long it takes DNA to mutate, and the average generation time of 25 years, researchers extrapolated that the cohanim Y chromosome pattern originated 2,100 to 3,250 years ago—which includes the time when Moses lived. According to religious documents, Moses' brother Aaron was the first priest.

The Jewish priest DNA signature also appears today among the Lemba, a population of South Africans with black skin. Researchers looked at them for the telltale gene variants because their customs suggest a Jewish origin—they do not eat pork (or hippopotamus), they circumcise their newborn sons, and they celebrate a weekly day of rest (**figure 1.7**).

To understand the extent and nuances of human genetic variation today, as well as to trace our “deep ancestries,” many people will need to have their genomes analyzed—not just members of illustrious families. This effort is gathering momentum. The Human Variome Project, for example, was planned in 1994 to catalog single genes, but the project now looks at SNPs across the genome, using their patterns to correlate genotypes to phenotypes that affect health.

An effort that is genealogical in focus is the Genographic Project. Begun with indigenous peoples, anyone can now send in a DNA sample for tracing the maternal and/or the paternal line back, possibly as far as about 56,000 years ago, when

the first modern humans left Africa and left descendants. Data from hundreds of thousands of people are being databased anonymously, and



Figure 1.7 Y chromosome DNA sequences reveal origins. The Lemba, a modern people with dark skin, have the same Y chromosome DNA sequences as the cohanim, a group of Jewish priests. The Lemba practiced Judaism long before DNA analysis became available.

compared to deduce likely migratory routes within and out of Africa. **Reading 16.2**, Should You Take a Genetic Ancestry Test?, provides details.

Health Care

Looking at diseases from a genetic point of view is changing health care. Many diseases, not just inherited ones, are now viewed as the consequence of complex interactions among genes and environmental factors. Even the classic single-gene diseases are sensitive to the environment. A child with cystic fibrosis (MIM 219700), for example, is more likely to suffer frequent respiratory infections if she regularly breathes second-hand smoke. A genetic approach to health is as much common sense as it is technological.

Diseases can result from altered proteins or too little or too much of a protein, or proteins made at the wrong place or time. Gene expression profiling studies are revealing the sets of genes that are turned on and off in specific cells and tissues as health declines. Genes also affect how people respond to particular drugs. For example, inheriting certain gene variants can make a person's body very slow at breaking down an anti-clotting drug, or extra sensitive to the drug. Such an individual could experience dangerous bleeding at the same dose that most patients tolerate. Identifying individual drug reactions based on genetics is a growing field called pharmacogenomics. **Table 1.3** lists some examples.

Single-Gene Diseases

Inherited illness caused by a single gene differs from other types of illnesses in several ways (**table 1.4**). In families, we can predict inheritance of a disease by knowing exactly how a person is related to an affected relative, discussed in chapter 4. In contrast, an infectious disease requires that a pathogen pass from one person to another, which is a much less predictable circumstance.

A second distinction of single-gene disorders is that the risk of developing symptoms can sometimes be predicted. This is possible because all cells harbor the mutation. A person with a family history of Huntington disease (HD; MIM 143100), for example, can have a blood test that detects the mutation at any age, even though symptoms typically do not occur until near age 40. *Bioethics: Choices for the Future* in chapter 4 discusses this further. Inheriting the HD mutation predicts illness with near certainty. For many conditions, predictive power is much

Table 1.3 Pharmacogenomic Tests

Antidepressants
Chemotherapies
HIV drugs
Smoking cessation drugs
Statins (cholesterol-lowering drugs)
Warfarin (anti-clotting)

Table 1.4 How Single-Gene Diseases Differ from Other Diseases

1. Risk can be predicted for family members.
2. Predictive (presymptomatic) testing may be possible.
3. Different populations may have different characteristic disease frequencies.
4. Correction of the underlying genetic abnormality may be possible.

lower. For example, inheriting one copy of a particular variant of a gene called *APOE* raises risk of developing Alzheimer disease by three-fold, and inheriting two copies raises it 15-fold. But without absolute risk estimates and no treatments for this disease, would you want to know?

A third feature of single-gene diseases is that they may be much more common in some populations than others. Genes do not like or dislike certain types of people; rather, mutations stay in certain populations because we marry people like ourselves. While it might not seem politically correct to offer a “Jewish genetic disease” screen, it makes biological and economic

sense—two dozen disorders are much more common in this population. A fourth characteristic of a genetic disease is that it may be “fixable” by altering the abnormal instructions.

Redefining Disease to Reflect Gene Expression

Diseases are increasingly being described in terms of gene expression patterns, which is not the same as detecting mutations. Gene expression refers to whether a gene is “turned on” or “turned off” from being transcribed and translated into protein (see Reading 1.1).

Tracking gene expression can reveal new information about diseases and show how diseases are related to each other. **Figure 1.8** shows part of a huge depiction of genetic disease called the “diseasome.” It connects diseases that share genes that show altered expression. Like most semantic webs that connect information from databases, the diseasome reveals relationships among diseases that were not obvious from traditional medical science, which is based on observing symptoms, detecting pathogens or parasites, or measuring changes in body fluid composition.

Some of the links and clusters in the diseasome are well-known, such as obesity, hypertension, and diabetes. Others are

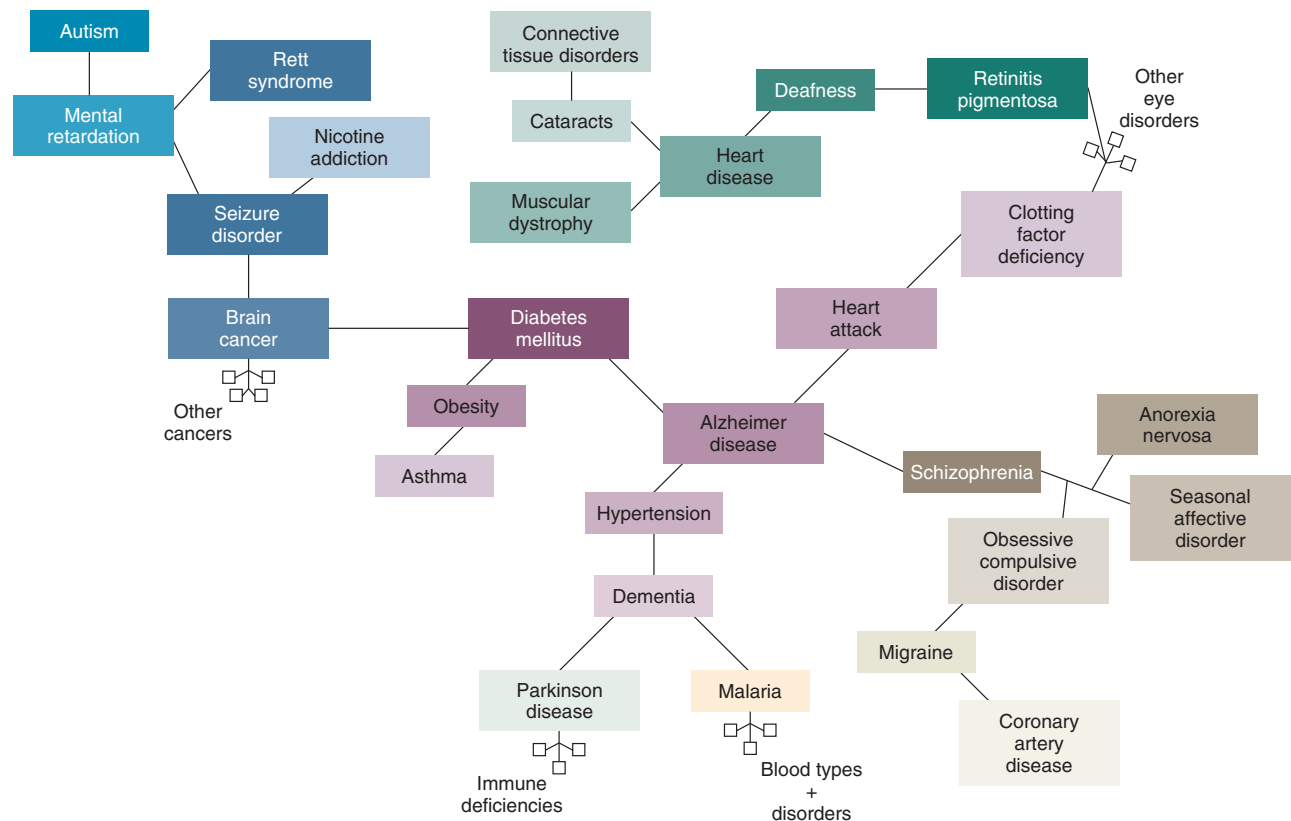


Figure 1.8 Part of the diseasome. This tool links diseases by shared gene expression. That is, a particular gene may be consistently overexpressed or underexpressed in two diseases, compared to the healthy condition. The lines refer to at least one gene connecting the disorders depicted in the squares. The conditions are not necessarily inherited because gene expression changes in all situations. The diseasome is an oversimplification in several ways. The same symptoms may have different causes, and each condition is associated with expression changes in more than one gene. Shading indicates conditions that may share a symptom. (Based on the work of A-L Barabási and colleagues.)

surprises, such as Duchenne muscular dystrophy (DMD; see figure 2.1) and heart attacks. The muscle disorder has no treatment, but heart attack does—researchers are now testing cardiac drugs on boys with DMD. In other cases, the association of a disease with genes whose expression goes up or down can suggest targets for new drugs.

The disease approach to defining and classifying diseases by their genetic underpinnings will have many practical consequences. It might alter the codes for different medical conditions, used in hospitals and for insurance. The World Health Organization may have to re-examine its lists of causes of death. Diseases with different symptoms might be found to be variations of the same underlying defect, whereas some conditions with similar symptoms might be found to be distinct at the molecular level.

Genetic Testing

Tests to identify about 1,200 single-gene disorders, most of them very rare, have been available for years. Direct-to-consumer (DTC) genetic testing, via websites and cheek cell samples, is bringing many kinds of DNA-based tests to many more people. Before passage of the Genetic Information Non-discrimination Act (GINA) in the United States in 2008, it was common for people to avoid genetic testing for fear of the misuse of genetic information or to take tests under false names so the result would not appear in their medical records. Some people refused to participate in clinical trials of new treatments if genetic information could be traced to them.

Under GINA, employers cannot use genetic information to hire, fire, or promote an employee, or require genetic testing. Similarly, health insurers cannot require genetic tests nor use the results to deny coverage. GINA also clearly defines a genetic test: It is an analysis of human DNA, RNA, chromosomes, proteins, or metabolites, to detect genotypes, mutations, or chromosomal changes. The law defines “genetic information” as tests or phenotypes (traits or symptoms) in individuals and/or families.

The long-awaited GINA legislation, however, raises new issues. Consider two patients with breast cancer—one with a strong family history and a known mutation, the other diagnosed after a routine mammogram, with no family history or identified mutation. A health insurer could refuse to cover the second woman, but not the first. Other limitations of GINA are that it does not apply to companies with fewer than 15 employees, it does not overrule state law, it does not protect privacy, and it does not spell out how discrimination will be punished. These concerns will be addressed as the law is put into practice.

In the long term, genetic tests, whether for single-gene disorders or the more common ones with associated genetic risks, may actually lower health care costs. If people know their inherited risks, they can forestall or ease symptoms that environmental factors might trigger—such as by eating healthy foods suited to their family history, not smoking, exercising regularly, avoiding risky behaviors, having frequent medical exams and screening tests, and beginning treatments earlier,

when they are more likely to be effective. The protection of GINA will also help recruit participants for clinical trials.

Treatments

Only a few single-gene diseases can be treated. Supplying a missing protein directly can prevent some symptoms, such as giving a clotting factor to a person with a bleeding disorder. Some inborn errors of metabolism (see Reading 2.1) in which an enzyme deficiency leads to build-up of a biochemical in cells, can be counteracted by tweaking diet to minimize the accumulation. Treatment at the DNA level—gene therapy—replaces the faulty instructions for producing the protein in cells that are affected in the illness.

For some genetic diseases, better understanding of how mutations cause the symptoms suggests that an existing drug for another condition might work. For example, experiments in mice with tuberous sclerosis complex, a disease that causes autism, memory deficits, and mental retardation in humans (MIM 191100), led to clinical trials of a drug, rapamycin, already in use to lessen transplant rejection. Tuberous sclerosis affects the same enzyme that the drug targets. Chapter 20 discusses various approaches to treating genetic disease.

Genome information is useful for treating infectious diseases, because the microorganisms and viruses that make us sick also have genetic material that can be sequenced and detected. In one interesting case, three patients died from infection 6 weeks after receiving organs from the same donor. All tests for known viruses and bacteria were negative, so medical researchers sampled DNA from the infected organs, removed human DNA sequences and those of known pathogens, and examined the remainder for sequences that resemble those of bacteria and viruses. This approach picked up genetic material from pathogens that cannot be grown in the laboratory. Using the DNA sequence information to deduce and reconstruct physical features of the pathogens, the researchers were able to identify a virus that caused the transplant recipients' deaths. Researchers then developed a diagnostic test for future transplant recipients who have the same symptoms.

Agriculture

The field of genetics arose from agriculture. Traditional agriculture is the controlled breeding of plants and animals to select individuals with certain combinations of inherited traits that are useful to us, such as seedless fruits or lean meat. **Biotechnology**, which is the use of organisms to produce goods (including foods and drugs) or services, is an outgrowth of agriculture.

One ancient example of biotechnology is using microorganisms to ferment fruits to manufacture alcoholic beverages, a technique the Babylonians used by 6000 B.C. Beer brewers in those days experimented with different yeast strains cultured under different conditions to control aroma, flavor, and color. Today, researchers have sequenced the genomes of the two types of yeast that are crossed to ferment lager beer, which requires lower temperatures than does ale. The work has shown that beers from different breweries around the world

have unique patterns of gene expression, suggesting ways to brew new types of beer.

Traditional agriculture is imprecise because it shuffles many genes—and, therefore, many traits—at a time, judging them by taste or appearance. In contrast, DNA-based techniques enable researchers to manipulate one gene at a time, adding control and precision to what is possible with traditional agriculture. Organisms altered to have new genes or to over- or underexpress their own genes are termed “genetically modified” (GM). If the organism has genes from another species, it is termed transgenic. Golden rice, for example, manufactures twenty-three times as much beta carotene (a vitamin A precursor) as unaltered rice. It has “transgenes” from corn and bacteria. Golden rice also stores twice as much iron as unaltered rice because one of its own genes is overexpressed. These nutritional boosts bred into edible rice strains may help prevent vitamin A and iron deficiencies in people who eat them.

People in the United States have been safely eating GM foods for more than a decade. In Europe, many people object to GM foods, on ethical grounds or based on fear. Officials in France and Austria have called such crops “not natural,” “corrupt,” and “heretical.” Food labels in Europe, and some in the United States, indicate whether a product is “GM-free.” Labeling foods can prevent allergic reaction to an ingredient in a food that wouldn’t naturally be there, such as a peanut protein in corn.

Field tests may not adequately predict the effects of GM crops on ecosystems. GM plants have been found far beyond where they were planted, thanks to wind pollination. Planting GM crops may also lead to extreme genetic uniformity, which could be disastrous. Some GM organisms, such as fish that grow to twice normal size or can survive at temperature extremes, may be so unusual that they disrupt ecosystems. **Figure 1.9** shows an artist’s rendition of these fears.

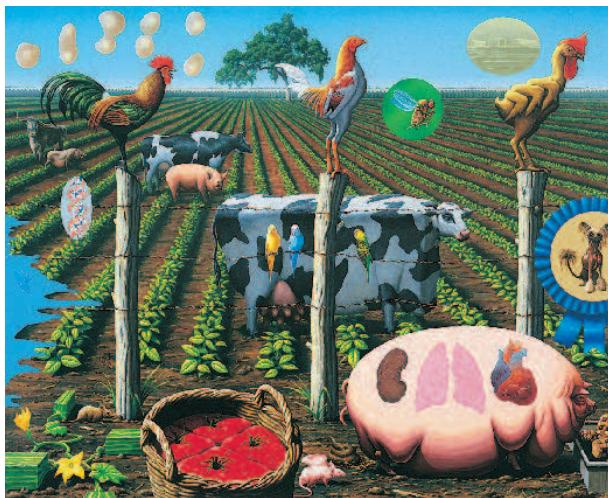


Figure 1.9 An artist’s view of biotechnology. Artist Alexis Rockman vividly captures some fears of biotechnology, including a pig used to incubate spare parts for sick humans, a muscle-boostered boxy cow, a featherless chicken with extra wings, a mini-warthog, and a mouse with a human ear growing out of its back.

Ecology

We share the planet with many thousands of other species. We aren’t familiar with many of Earth’s residents because we can’t observe their habitats, or we can’t grow them in laboratories. “Metagenomics” is a field that is revealing and describing much of the invisible living world by sequencing all of the DNA in a particular habitat. Such areas range from soil, to an insect’s gut, to garbage. Metagenomics studies are revealing how species interact, and may yield new drugs and reveal novel energy sources.

Metagenomics researchers collect and sequence DNA and consult databases of known genes and genomes to imagine what the organisms might be like. One of the first metagenomics projects described life in the Sargasso Sea. This 2-million-square-mile oval area off the coast of Bermuda has long been thought to lack life beneath its thick cover of seaweed, which is so abundant that Christopher Columbus thought he’d reached land when his ships came upon it. Many a vessel has been lost in the Sargasso Sea, which includes the area known as the Bermuda Triangle. When researchers sampled the depths, they collected more than a billion DNA bases, representing about 1,800 microbial species, including at least 148 not seen before. More than a million new genes were discovered.

A favorite site for metagenomics analysis is the human body. The Human Microbiome Project is exploring the other forms of life within us. Genome profiling on various parts of our anatomy reveals that 90 percent of the cells in a human body are not actually human! A human body is, in fact, a vast ecosystem. This is possible because bacterial cells are so much smaller than ours. Humans have a “core microbiome” of bacterial species that everyone has, but also many others that reflect our differing environments, habits, ages, diets, and health.

Most of our bacterial residents live in our digestive tracts—about 10 trillion of them. The human mouth is home to about 500 different species of bacteria, only about 150 of which can grow in the laboratory. Analysis of their genomes yields practical information. For example, the genome of one bacterium, *Treponema denticola*, showed how it survives amid the films other bacteria form in the mouth, and how it causes gum disease. Sequencing genes in saliva from people from all over the world reveals that we are just as different in this regard from our neighbors as from people on the other side of the globe.

The other end of the digestive tract is easy to study too, because feces are very accessible research materials that are chock-full of bacteria from the intestines. One study examined soiled diapers from babies regularly during their first year, chronicling the establishment of the gut bacterial community. Newborns start out with blank slates—clean intestines—and after various bacteria come and go, very similar species remain in all the children by their first birthdays. Researchers study the bacteria that live between our mouths and anuses by looking at people who receive intestinal transplants, a very rare procedure. Intestines that are transplanted are first flushed clean of the donor’s bacteria. Researchers can sample bacteria through an opening made in the abdominal wall of the recipient. The few willing participants so far reveal that people are unique in

Bioethics: Choices for the Future

Genetic Testing and Privacy

The field of bioethics began in the 1950s and 1960s as a branch of philosophy that addressed issues raised by medical experimentation during World War II. Bioethics initially centered on matters of informed consent, paternalism, autonomy, allocation of scarce medical resources, justice, and definitions of life and death. Today, the field covers medical and biotechnologies and the choices and dilemmas they present. Genetic testing is at the forefront of twenty-first-century bioethics because its informational nature affects privacy. Consider these situations.

Testing Tissue from Deceased Children

When parents approve genetic testing for a sick child, they usually assume that their consent applies only when the child is still living, but research may continue after the child is gone. If a newly discovered gene function explains the condition of a child who had never received an accurate diagnosis, should the parents be informed? Would doing so reopen wounds, or provide helpful information?

The consensus of medical and scientific organizations is that posthumous genetic test information should be disclosed only if the results have been validated (confirmed), the results can lead to testing or treatment for others, and if the parents have not indicated that they do not want to know. For example, several years after a 7-year-old girl died of then-mysterious symptoms, her mother read an article about Rett syndrome (MIM 312750), and thought it described her daughter. Girls with Rett syndrome (boys are not affected) have small head, hands, and feet; poor socialization skills; cognitive impairment; and a characteristic repetitive movement (hand-wringing). They may be unable to otherwise move, and have seizures or digestive problems. Researchers confirmed the mother's suspicions by testing DNA extracted from a baby tooth she had saved. Finally having a diagnosis made it possible to test the other children in the family, who were not affected and could therefore not pass on the disease. Considering the current pace of gene discovery, it is likely that more posthumous genetic tests will be done in the future.

The Military

A new recruit hopes that the DNA sample that he or she gives when military service begins is never used—it is stored so that remains can be identified. Up until now, genetic tests have only been performed for two specific illnesses that could affect soldiers under certain environmental conditions. Carriers of sickle cell disease (MIM 603903) can develop painful blocked circulation at high altitudes, and carriers of G6PD deficiency (MIM 305900) react badly to anti-malaria medication. Carriers wear red bands on their arms to alert officers to be certain that they avoid the environments that could harm them.

The passage of GINA (the Genetic Information Nondiscrimination Act) has led to more precise definitions of genetic disease in the military, even though the law does not apply specifically to the armed forces. In the past, in determining benefits, the military assumed that any illness present when a soldier left military service that was not noted on entry was caused by serving, “with the exception of congenital and hereditary conditions.” Such wording discouraged genetic testing, because test results indicating future disease would be interpreted to mean a pre-existing condition. This is no longer the case. The National Defense Authorization Act of 2008 makes it clear that detecting a disease-causing gene mutation before symptoms begin does not constitute a medical diagnosis, and therefore cannot be used as a reason to deny benefits.

In the future, the military may use genetic information to identify soldiers at risk for such conditions as depression and post-traumatic stress disorder. Deployments can be tailored to risks, minimizing suffering.

Genome-Wide Association Studies and Disappearing Privacy

The first genome-wide association studies typed people for only a few hundred SNPs. This limited analysis ensured privacy because there were many more people than genotypes, so that it was highly unlikely that an individual could be identified by being the only one to have a particular genotype. That is no longer true. As studies now probe a million or more SNPs, an algorithm can analyze study data and match an individual to a genotype and trace that genotype to a particular group being investigated—revealing, for example, that a person has a particular disease. That is, the more ways that we can detect that people vary, the easier it is to identify any one of them. It is a little like adding four digits to a zip code, or more area codes to phone numbers, to increase the pool of identifiers. Several government DNA databases pulled their data from open access once an astute researcher discovered the transparency.

Questions for Discussion

1. What should be included in an informed consent document that would sensitively ask parents if they would like to receive research updates on their child's inherited disease after the child has passed away?
2. If a genetic test on a sick child, person in the military, or participant in a clinical trial or other experiment reveals a mutation that could harm a blood relative, should the first person's privacy be sacrificed to inform the second person?
3. What measures can physicians, the military, and researchers take to ensure that privacy of genetic information is maintained?

their “gut microbiome,” but that those whose bacterial species stay about the same over time are healthier than those whose bacterial types fluctuate.

In parallel to metagenomics, several projects are exploring biodiversity with DNA tags to “bar-code” species, rather than sequencing entire genomes. DNA sequences that vary reveal more about ancestries, because they are informational, than do comparisons of physical features, such as body shape or size, which formed the basis of traditional taxonomy (biological classification).

A Global Perspective

Because genetics so intimately affects us, it cannot be considered solely as a branch of life science. Equal access to testing, misuse of information, and abuse of genetics to intentionally cause harm are compelling issues that parallel scientific progress.

Genetics and genomics are spawning technologies that may vastly improve quality of life. But at first, tests and treatments will be costly and not widely available. While advantaged people in economically and politically stable nations may look forward to genome-based individualized health care, poor people in other nations just try to survive, often lacking basic vaccines and medicines. In an African nation where two out of five children suffer from AIDS and many die from other infectious diseases, newborn screening for rare single-gene defects hardly seems practical. However, genetic disorders weaken people so that they become more susceptible to infectious diseases, which they can pass to others.

Human genome information can ultimately benefit everyone. Genome information from humans and our pathogens and parasites is revealing new drug targets. Global organizations, including the United Nations, World Health Organization, and the World Bank, are discussing how nations can share new diagnostic tests and therapeutics that arise from genome information.

Individual nations are adopting approaches that exploit their particular strengths (table 1.5). India, for example, has many highly inbred populations with excellent genealogical records, and is home to one-fifth of the world’s population. Studies of genetic variation in East Africa are especially important because this region is the cradle of humanity—home of our forebears. The human genome belongs to us all, but efforts from around the world will tell us what our differences are and how they arose. *Bioethics: Choices for the Future* discusses instances when genetic testing can be intrusive.

Table 1.5 Nations Plan for Genomic Medicine

Nation	Program
China	The genomes of 100 people are being sequenced.
Gambia	A DNA databank has samples from 57,000 people.
India	A national databank stores DNA from 15,000 people. A company is genotyping the entire Parsi population of 69,000. Other efforts are examining why many drugs only help some people. Laws prevent foreign researchers from sampling tissue from Indians without permission.
Mexico	The National Institute for Genomic Medicine has genotyped 1,200+ people to look for correlations to common diseases. “Safari research” legislation requires approval for foreign researchers to sample DNA from Mexicans.
South Africa	Studies of human genetic diversity among indigenous tribes and susceptibility to HIV and tuberculosis among many populations are underway.
Thailand	A database stores information on genetic susceptibility to dengue fever, malaria, other infectious diseases, and posttraumatic stress disorder from the 2004 tsunami.

Key Concepts

1. Genetics has diverse applications. Matching DNA sequences can clarify relationships, which is useful in forensics, establishing identity, and understanding historical events.
2. Inherited disease differs from other disorders in its predictability; characteristic frequencies in different populations; and the potential of gene therapy.
3. Agriculture and biotechnology apply genetic principles.
4. Collecting DNA from habitats and identifying the sequences in databases is a new way to analyze ecosystems.
5. Human genome information has tremendous potential but must be carefully managed.

Summary

1.1 Introducing Genes

1. **Genes** are the instructions to manufacture proteins, which determine inherited traits.
2. A **genome** is a complete set of genetic information. A **cell**, the unit of life, contains two genomes of **DNA**. **Genomics** is the study of many genes and their interactions.

1.2 Levels of Genetics

3. Genes encode proteins and the **RNA** molecules that synthesize proteins. RNA carries the gene sequence information so that it can be utilized, while the DNA is transmitted when the cell divides. Much of the genome does not encode protein.

- Variants of a gene, called **alleles**, arise by **mutation**. Alleles may differ slightly from one another, but encode the same product. A polymorphism is a site or sequence of DNA that varies in one percent or more of a population.
- Genome-wide association studies** compare landmarks across the genomes among individuals who share a trait. **Gene expression profiling** examines which genes are more or less active in particular cell types.
- Chromosomes** consist of DNA and protein. The 22 types of **autosomes** do not include genes that specify sex. The X and Y **sex chromosomes** bear genes that determine sex.
- Cells **differentiate** by expressing subsets of genes. **Stem cells** divide to yield other stem cells and cells that differentiate.
- The **phenotype** is the gene's expression. An allele combination constitutes the **genotype**. Alleles may be **dominant** (exerting an effect in a single copy) or **recessive** (requiring two copies for expression).
- Pedigrees are diagrams used to study traits in families.
- Genetic populations are defined by their collections of alleles, termed the **gene pool**. Genome comparisons among species reveal evolutionary relationships.

1.3 Genes and Their Environment

- Single genes determine Mendelian traits. **Multifactorial** traits reflect the influence of one or more genes and the environment. Recurrence of a Mendelian trait is predicted based on Mendel's laws; predicting the recurrence of a multifactorial trait is more difficult.
- Genetic determinism** is the idea that the expression of an inherited trait cannot be changed.

1.4 Applications of Genetics

- DNA profiling can establish identity, relationships, and origins.
- In health care, single-gene diseases are more predictable than other diseases, but gene expression profiling is revealing how many types of diseases are related.
- Agriculture is selective breeding. **Biotechnology** is the use of organisms or their parts for human purposes. A transgenic organism harbors a gene or genes from a different species.
- In metagenomics, DNA collected from habitats, including the human body, is used to reconstruct ecosystems.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

- Place the following terms in size order, from largest to smallest, based on the structures or concepts they represent:
 - chromosome
 - gene pool
 - gene
 - DNA
 - genome
- Distinguish between:
 - an autosome and a sex chromosome
 - genotype and phenotype
 - DNA and RNA
 - recessive and dominant traits
 - pedigrees and karyotypes
 - gene and genome
- Explain how DNA encodes information.
- Explain how all humans have the same genes, but vary genetically.
- Explain how a genome-wide association study, gene expression profiling, and DNA sequencing of a gene or genome differ.
- Explain how all cells in a person's body have the same genome, but are of hundreds of different types that look and function differently.
- Suggest a practical example of gene expression profiling.
- Explain the protections under the Genetic Information Nondiscrimination Act, and the limitations.
- Explain what an application of a "diseasome" type of map, such as in figure 1.8, might provide.
- Cite an example of a phrase that illustrates genetic determinism.
- Give an example of a genome that is in a human body, but is not human.

Applied Questions

- If you were ordering a genetic test panel, which traits and health risks would you like to know about, and why?
- Two roommates go grocery shopping and purchase several packages of cookies that supposedly each provide 100 calories. After a semester of eating the snacks, one roommate has gained 6 pounds, but the other hasn't. Assuming that other dietary and exercise habits are similar, explain the roommates' different response to the cookies.

3. A study comparing feces of vegetarians, people who eat mostly meat (carnivores), and people who eat a variety of foods (omnivores) found that the microbiome of the vegetarians is much more diverse than that of the other types of diners. Explain why this might be so.
4. One variant in the DNA sequence for the gene that encodes part of the oxygen-carrying blood protein hemoglobin differs in people who have sickle cell disease. Newborns are tested for this mutation. Is this a single-gene test, a genome sequencing, a genome-wide association study, or a gene expression profile?
5. Consider the following two studies:
 - Gout is a form of arthritis that often begins with pain in the big toe. In one study, researchers looked at 500,000 SNPs in 100 people with gout and 100 who do not have gout, and found a very distinctive pattern in the people with painful toes.
 - About 1 percent of people who take cholesterol-lowering drugs (statins) experience muscle pain. Researchers discovered that their muscle cells have different numbers and types of mRNA molecules than the majority of people who tolerate the drugs well.
6. Which description is of a genome-wide association study and which a gene expression study?
6. A 54-year-old man is turned down for life insurance because testing following a heart attack revealed that he had inherited cardiac myopathy, and this had most likely caused the attack. He cites GINA, but the insurer says that the law does not apply to his case. Who is correct?
7. How will GINA benefit
 - a. health care consumers?
 - b. employers?
 - c. insurers?
 - d. researchers?
8. An ad for a skin cream proclaims it will “boost genes’ activity and stimulate the production of youth proteins.” Which technology described in the chapter could be used to test the ad’s claim?

Web Activities

9. Consult a website for a direct-to-consumer genetic testing company, such as 23andMe, Navigenics, or deCODE Genetics. Choose three tests, and explain why you would want to take them. Also discuss a genetic test that you would not wish to take, and explain why not.
10. Many organizations are using DNA bar codes to classify species. Consult the websites for one of the following organizations and describe an example of how they are using DNA sequences:
 - Consortium for the Barcode of Life (International)
 - Canadian Barcode of Life Network
 - Species 2000 (UK)
 - Encyclopedia of Life (Wikipedia)
11. Human microbiome projects have different goals. Consult the websites for two of the following projects and compare their approaches:
 - The Human Microbiome Project (NIH)
 - Meta-Gut (China)
 - Metagenomics of the Human Intestinal Tract (European Commission)
 - Human Gastric Microbiome (Singapore)
 - Australian Urogenital Microbiome Consortium
 - Human MetaGenome Consortium (Japan)
 - Canadian Microbiome Initiative
12. Look at the website for the McLaughlin-Rotman Centre for Global Health (www.mrcglobal.org). Describe a nation’s plan to embrace genomic medicine.

Forensics Focus

13. Consult the websites for a television program that uses or is based on forensics (*CSI* or *Law and Order*, for example), and find an episode in which species other than humans are critical to the case. Explain how DNA bar coding could help to solve the crime.
14. On an episode of the television program *House*, the main character, Dr. House, knew from age 12 that his biological father was a family friend, not the man who raised him. At his supposed father’s funeral, the good doctor knelt over the body in the casket and sneakily snipped a bit of skin from the corpse’s earlobe—for a DNA test.
 - a. Do you think that this action was an invasion of anyone’s privacy? Was Dr. House justified?
 - b. Dr. House often orders treatments for patients based on observing symptoms. Suggest a way that he can use DNA testing to refine his diagnoses.



When Michael M. received stem cells to heal his eyes, his sight (sensation of light) was restored, but not his vision (his brain's perception of the images). Slowly, his brain caught up with his senses, and he was able to see his family for the first time.

CHAPTER

2

Cells

Chapter Contents

- 2.1 Introducing Cells**
- 2.2 Cell Components**
 - Chemical Constituents
 - Organelles
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 - The Cytoskeleton
- 2.3 Cell Division and Death**
 - The Cell Cycle
 - Apoptosis
- 2.4 Cell-Cell Interactions**
 - Signal Transduction
 - Cellular Adhesion
- 2.5 Stem Cells**
 - Cell Lineages
 - Stem Cells in Health Care

Stem Cells Restore Sight, But Not Vision

In 1960, 3-year-old Michael M. lost his left eye in an accident. Because much of the vision in his right eye was already impaired from scars on the cornea (the transparent outer layer) he could see only distant, dim light. Several corneal transplants failed, adding more scar tissue. At age 39, Michael received stem cells from a donated cornea and the tissue finally regrew. Researchers learned just recently that corneal transplants work only if the transplanted tissue includes stem cells.

After the transplant, Michael could see his wife and two sons for the first time. But he quickly learned that vision is more than seeing—his brain had to interpret images. Because the development of his visual system had stalled, and he had only one eye, he could discern shapes and colors, but not three-dimensional objects, such as facial details. In fact, he had been more comfortable skiing blind, using verbal cues, than he was with sight—the looming trees were terrifying. It took years for Michael's brain to catch up to his rejuvenated eye.

The eye actually contains several varieties of stem cells, and they may be useful to heal more than visual illnesses and injuries. A single layer of cells called the retinal pigment epithelium, for example, forms at the back of the eye in an embryo, where it replenishes cells of the retina. These cells are typically discarded during eye surgery, but when cultured in a dish with a “cocktail” used for stem cells, can become nearly any cell type. One day, it might be possible to treat a brain disease, such as Parkinson disease, using a patient's own eye stem cells—without sacrificing vision.

2.1 Introducing Cells

The activities and abnormalities of cells underlie our inherited traits, quirks, and illnesses. Understanding cell function reveals how a healthy body works, and how it develops from one cell to trillions. Understanding what goes wrong in certain cells to cause pain or other symptoms can suggest ways to treat the condition—we learn what must be repaired or replaced. In Duchenne muscular dystrophy (MIM 310200), for example, the reason that a little boy's calf muscles are overdeveloped is that he cannot stand normally because other muscles are weak. The affected cells lack a protein that supports the cells' shape during forceful contractions (**figure 2.1**). Identifying the protein revealed exactly what must be replaced—but doing so has been difficult because many muscle cells must be corrected.

Our bodies include more than 260 variations on the cellular theme. Differentiated cell types include bone and blood, nerve and muscle, and subtypes of those. These are **somatic cells**, also called body cells. Somatic cells have two copies of the genome and are said to be **diploid**. In contrast, the rarer sperm and egg have one copy of the genome and are **haploid**. The meeting of sperm and egg restores the diploid state. Especially important in many-celled organisms are **stem cells**, which are diploid cells that both give rise to differentiated cells and replicate themselves, a characteristic called self-renewal. Stem cells enable a body to develop, grow, and repair damage.

Cells interact. They send, receive, and respond to information. Some cells aggregate with others of like function, forming tissues, which in turn interact to form organs and organ systems. Other cells move about the body. Cell numbers are important, too—they are critical to development, growth, and healing. Staying healthy reflects a precise balance between cell division, which adds cells, and cell death, which takes them away.

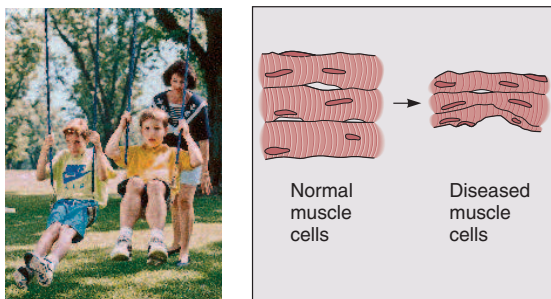


Figure 2.1 Genetic disease at the whole-person and cellular levels. An early sign of the boy on the right's Duchenne muscular dystrophy is overdeveloped calf muscles that result from his inability to rise from a sitting position the usual way. Lack of the protein dystrophin causes his skeletal muscle cells to collapse when they contract. Stem cells can treat this condition in mice and dogs.

2.2 Cell Components

All cells share certain features that enable them to perform the basic life functions of reproduction, growth, response to stimuli, and energy use. Specialized features emerge as cells express different subsets of the thousands of protein-encoding genes. Many other genes control which protein-encoding genes a cell expresses.

Other multicellular organisms, including other animals, fungi, and plants, also have differentiated cells. Some single-celled organisms, such as the familiar paramecium and amoeba, have very distinctive cells as complex as our own. The most abundant organisms on the planet, however, are simpler and single-celled. These microorganisms are nonetheless successful life forms because they have occupied Earth much longer than we have, and even live in our bodies.

Biologists recognize three broad varieties of cells that define three major “domains” of life: the Archaea, the Bacteria, and the Eukarya. A domain is a broader classification than the familiar kingdom.

Members of the archaea and bacteria are single-celled, but they differ from each other in the sequences of many of their genetic molecules and in the types of molecules in their membranes. Archaea and bacteria are, however, both **prokaryotes**, which means that they lack a **nucleus**, the structure that contains DNA in the cells of other types of organisms.

The third domain of life, the Eukarya or **eukaryotes**, includes single-celled organisms that have nuclei, as well as all multicellular organisms such as ourselves (**figure 2.2**). Eukaryotic cells are also distinguished from prokaryotic cells in that they have structures called **organelles**, which perform specific functions. The cells of all three domains contain globular assemblies of RNA and protein called **ribosomes** that are

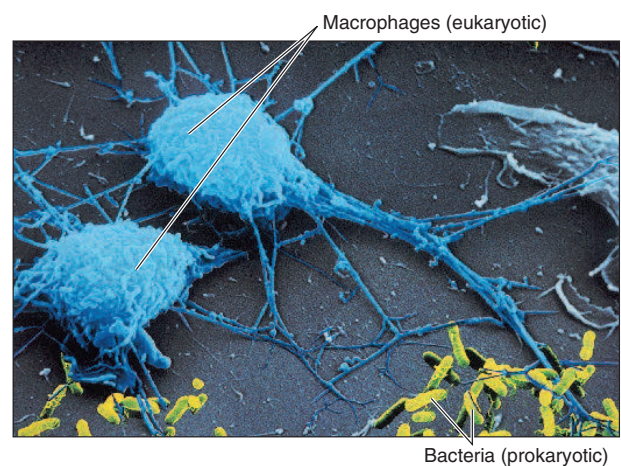


Figure 2.2 Eukaryotic and prokaryotic cells. A human cell is eukaryotic and much more complex than a bacterial cell, while an archaean cell looks much like a bacterial cell. Here, human macrophages (blue) capture bacteria (yellow). Note how much larger the human cells are. (A few types of giant bacteria are larger than some of the smaller human cell types.)

essential for protein synthesis. The eukaryotes may have arisen from an ancient fusion of a bacterium with an archaean.

Chemical Constituents

Cells are composed of molecules. Some of the chemicals of life (biochemicals) are so large that they are called macromolecules.

The major macromolecules that make up cells and are used by them as fuel are **carbohydrates** (sugars and starches), **lipids** (fats and oils), **proteins**, and **nucleic acids** (DNA and RNA). Cells require vitamins and minerals in much smaller amounts.

Carbohydrates provide energy and contribute to cell structure. Lipids form the basis of several types of hormones, form membranes, provide insulation, and store energy. Proteins have many diverse functions in the human body. They participate in blood clotting, nerve transmission, and muscle contraction and form the bulk of the body's connective tissue. Antibodies that fight bacterial infection are proteins. **Enzymes** are especially important proteins because they facilitate, or catalyze, biochemical reactions so that they occur swiftly enough to sustain life. Most important to the study of genetics are the nucleic acids DNA and RNA, which translate information from past generations into specific collections of proteins that give a cell its individual characteristics.

Macromolecules often combine in cells, forming larger structures. For example, the membranes that surround cells and compartmentalize their interiors consist of double layers (bilayers) of lipids embedded with carbohydrates, proteins, and other lipids.

Life is based on the chemical principles that govern all matter; genetics is based on a highly organized subset of the chemical reactions of life. **Reading 2.1** describes some drastic effects that result from major biochemical abnormalities.

Organelles

A typical eukaryotic cell holds a thousand times the volume of a bacterial or archaeal cell. To carry out the activities of life in such a large cell, organelles divide the labor by partitioning off certain areas or serving specific functions. The coordinated functioning of the organelles in a eukaryotic cell is much like the organization of departments in a big-box store, compared to the prokaryote-like simplicity of a small grocery store. In general, organelles keep related biochemicals and structures close enough to one another to interact efficiently. This eliminates the need to maintain a high concentration of a particular biochemical throughout the cell.

Organelles have a variety of functions. They enable a cell to retain as well as to use its genetic instructions, acquire energy, secrete substances, and dismantle debris. Saclike organelles sequester biochemicals that might harm other cellular constituents. Some organelles consist of membranes studded with enzymes embedded in the order in which they participate in the chemical reactions that produce a particular molecule. **Figure 2.3** depicts organelles.

The most prominent organelle of most cells is the nucleus. It is enclosed in a layer called the nuclear envelope. Nuclear pores are rings of proteins that allow certain biochemicals to exit or enter the nucleus (**figure 2.4**).

On the inner face of the nuclear membrane is a layer of fibrous material called the nuclear lamina. This layer has several important functions. The DNA within the nucleus touches the nuclear lamina as the cell divides. The nuclear lamina also provides mechanical support and holds in place the nuclear pores. Chapter 3 discusses very rare, accelerated aging disorders that result from an abnormal nuclear lamina.

Within the nucleus, an area that appears darkened under a microscope, called the nucleolus ("little nucleus"), is the site of ribosome production. The nucleus is filled with DNA complexed with many proteins to form chromosomes. Other proteins form fibers that give the nucleus a roughly spherical shape. RNA is abundant too, as are enzymes and proteins required to synthesize RNA from DNA. The fluid in the nucleus, minus these contents, is called nucleoplasm.

The remainder of the cell—that is, everything but the nucleus, organelles, and the outer boundary, or **plasma membrane**—is **cytoplasm**. Other cellular components include stored proteins, carbohydrates, and lipids; pigment molecules; and various other small chemicals. We now take a closer look at three cellular functions.

Secretion—The Eukaryotic Production Line

Organelles interact in ways that coordinate basic life functions and sculpt the characteristics of specialized cell types. Secretion, which is the release of a substance from a cell, illustrates how organelles function together.

Secretion begins when the body sends a biochemical message to a cell to begin producing a particular substance. For example, when a newborn first suckles the mother's breast, the stimulation causes her brain to release hormones that signal cells in her breast, called lactocytes, to rapidly increase the production of the complex mixture that makes up milk (**figure 2.5**). In response, information in certain genes is copied into molecules of **messenger RNA** (mRNA), which then exit the nucleus (see steps 1 and 2 in figure 2.5). In the cytoplasm, the mRNAs, with the help of ribosomes and another type of RNA called **transfer RNA**, direct the manufacture of milk proteins. These include nutritive proteins called caseins, antibodies that protect against infection, and enzymes.

Most protein synthesis occurs on a maze of interconnected membranous tubules and sacs called the **endoplasmic reticulum** (ER) (see step 3 in figure 2.5). The ER winds from the nuclear envelope outward to the plasma membrane. The section of ER nearest the nucleus, which is flattened and studded with ribosomes, is called rough ER, because the ribosomes make it appear fuzzy when viewed under an electron microscope. Messenger RNA attaches to the ribosomes on the rough ER. Amino acids from the cytoplasm are then linked, following the instructions in the mRNA's sequence, to form particular proteins that will either exit the cell or become part of membranes (step 3, figure 2.5). Proteins are also synthesized

Reading 2.1

Inborn Errors of Metabolism Affect the Major Biomolecules

Enzymes are proteins that speed specific chemical reactions, and, therefore, ultimately control a cell's production of all types of macromolecules. When the gene that encodes an enzyme mutates so that the enzyme is not produced or cannot function, the result can be too much or too little of the product of the biochemical reaction that the enzyme catalyzes. These biochemical buildups and breakdowns may cause symptoms. Genetic disorders that result from deficient or absent enzymes are called "inborn errors of metabolism." Following are some examples.

Carbohydrates

The newborn yelled and pulled up her chubby legs in pain a few hours after each feeding. She developed watery diarrhea, even though she was breastfed. Finally, a doctor diagnosed *lactase deficiency* (MIM 223000)—lack of the enzyme lactase, which enables the digestive system to break down the carbohydrate lactose. Bacteria multiplied in the undigested lactose in the child's intestines, producing gas, cramps, and bloating. Switching to a soybean-based, lactose-free infant formula helped. A different disorder with milder symptoms is lactose intolerance (MIM 150200), common in adults (see the opening essay to chapter 15).

Lipids

A sudden sharp pain began in the man's arm and spread to his chest. At age 36, he was younger than most people who suffer heart attacks, but he had inherited a gene variant that halved the number of protein receptors for cholesterol on his liver cells. Because cholesterol could not enter the liver cells efficiently, it built up in his arteries, constricting blood flow in his heart and eventually causing a mild heart attack. A fatty diet and lack of exercise had accelerated his *familial hypercholesterolemia*. A cholesterol-lowering drug and lifestyle changes lowered his risk of suffering future heart attacks.

Proteins

Newborn Tim slept most of the time, and he vomited so often that he hardly grew. A blood test revealed *maple syrup urine disease* (MIM 248600), so named because this inborn error of metabolism makes urine smell like maple syrup. Tim could not digest three types of amino acids (protein building blocks), which accumulated in his bloodstream. A diet very low in these amino acids controlled the symptoms. Today this inborn error is one of many dozen that are detected with blood tests shortly after birth. Newborn screening is discussed in chapter 20.

Nucleic Acids

From birth, Troy's wet diapers contained orange, sandlike particles, but otherwise he seemed healthy. By 6 months of age, he was in pain when urinating. A physician noted that Troy's writhing movements were involuntary rather than normal crawling.

The orange particles in Troy's diaper indicated *Lesch-Nyhan syndrome* (MIM 300322), caused by the deficiency of an enzyme called HGPRT. Troy's body could not recycle two of the four types of DNA building blocks, instead converting them into uric acid, which crystallizes in urine. Other symptoms that began later were not as easy to explain—severe mental retardation, seizures, and aggressive and self-destructive behavior. By age 3, he responded to stress by uncontrollably biting his fingers, lips, and shoulders. On doctors' advice, his parents had his teeth removed to keep him from harming himself, and he was kept in restraints. Troy would probably die before the age of 30 of kidney failure or infection.

Vitamins

Vitamins enable the body to use the carbohydrates, lipids, and proteins we eat. Julie inherited *biotinidase deficiency* (MIM 253260), which greatly slows her body's use of the vitamin biotin. If Julie hadn't been diagnosed as a newborn and quickly started on biotin supplements, by early childhood she would have shown biotin deficiency symptoms: mental retardation, seizures, skin rash, and loss of hearing, vision, and hair. Her slow growth, caused by her body's inability to extract energy from nutrients, would have eventually proved lethal.

Minerals

Ingrid, in her thirties, lived in the geriatric ward of a mental hospital, unable to talk or walk. She grinned and drooled, but she was alert and communicated using a computer. When she was a healthy high-school senior, symptoms of *Wilson disease* (MIM 277900) began as her weakened liver could no longer control the excess copper her digestive tract absorbed from food. The initial symptoms were stomachaches, headaches, and an inflamed liver (hepatitis). Then other changes began—slurred speech; loss of balance; a gravelly, low-pitched voice; and altered handwriting. A psychiatrist noted the telltale greenish rings around her irises, caused by copper buildup,

and diagnosed Wilson disease (**figure 1**). Finally Ingrid received penicillamine, which enabled her to excrete the excess copper in her urine. The treatment halted the course of the illness, saving her life. She now lives with a relative.



Figure 1 Wilson disease. A greenish ring around the brownish iris is one sign of the copper buildup of Wilson disease.

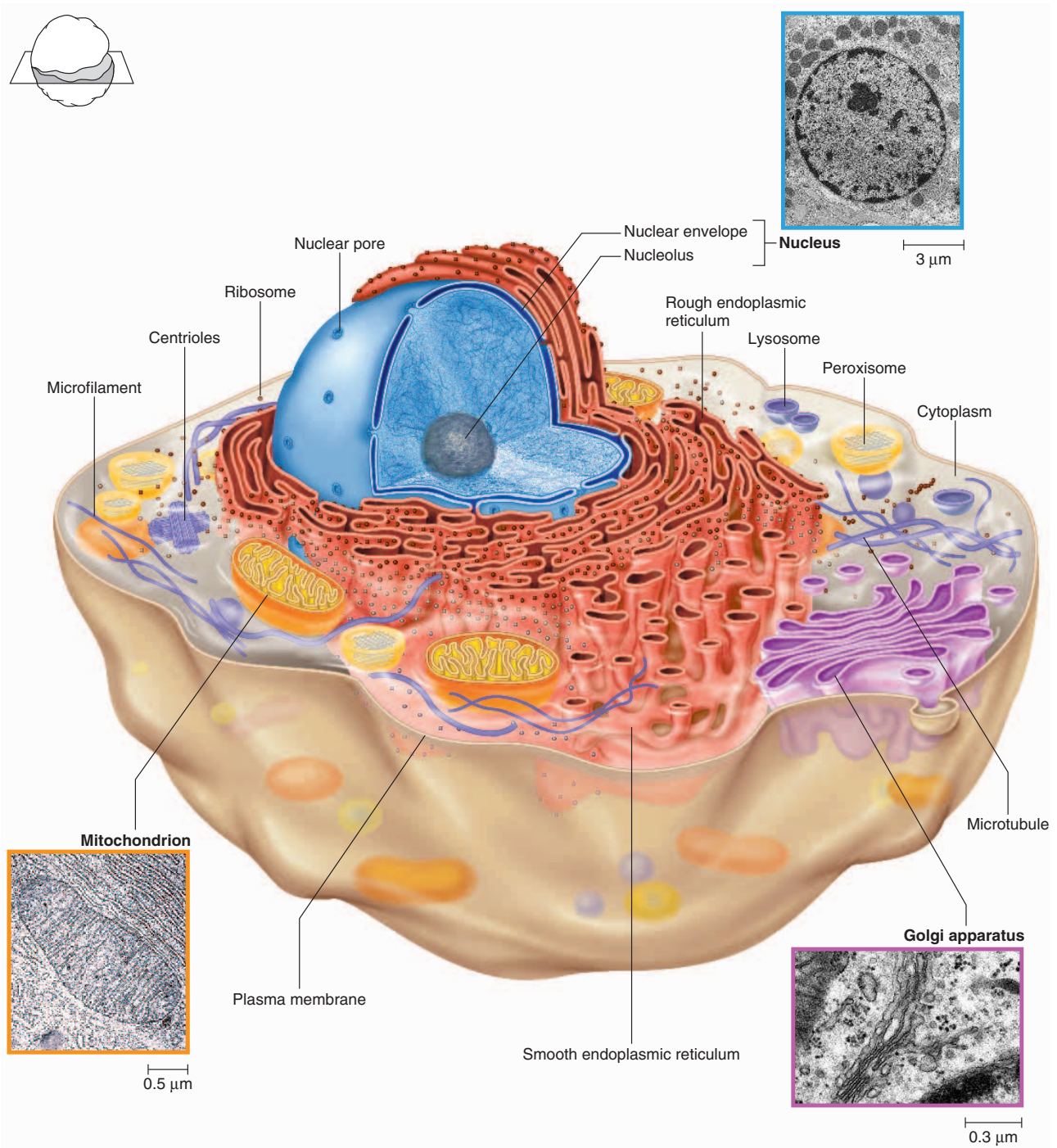


Figure 2.3 Generalized animal cell. Organelles provide specialized functions for the cell. Most of these structures are transparent; colors are used here to distinguish them. Different cell types have different numbers of organelles. All cell types have a single nucleus, except for red blood cells, which expel their nuclei as they mature.

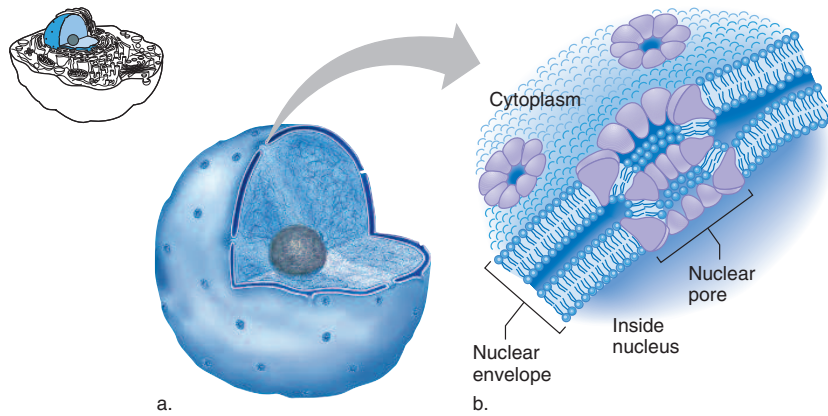


Figure 2.4 The nucleus is the genetic headquarters. (a) The largest structure in a typical human cell, the nucleus lies within two membrane layers that make up the nuclear envelope (b). Nuclear pores allow specific molecules to move in and out of the nucleus through the envelope.

on ribosomes not associated with the ER. These proteins remain in the cytoplasm.

The ER acts as a quality control center for the cell. Its chemical environment enables the forming protein to start folding into the three-dimensional shape necessary for its specific function. Misfolded proteins are pulled out of the ER and degraded, much as an obviously defective toy might be pulled from an assembly line at a toy factory and discarded. Misfolded proteins can cause disease, as discussed further in chapter 10.

As the rough ER winds out toward the plasma membrane, the ribosomes become fewer, and the tubules widen, forming a section called smooth ER. Here, lipids are made and added to the proteins arriving from the rough ER (step 4, figure 2.5). The lipids

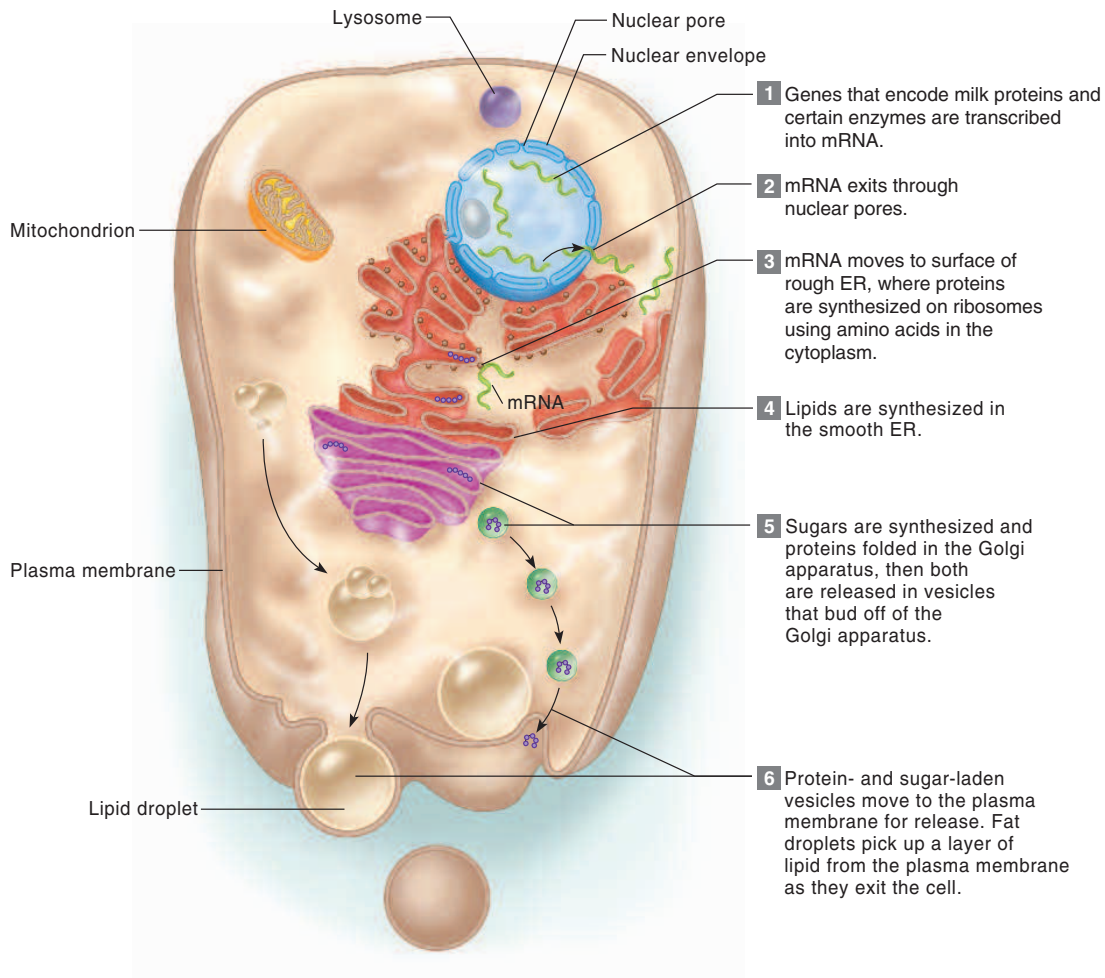


Figure 2.5 Secretion: making milk. Milk production and secretion illustrate organelle functions and interactions in a cell from a mammary gland: (1) through (6) indicate the order in which organelles participate in this process. Lipids are secreted in separate droplets from proteins and their attached sugars. This cell is highly simplified.

and proteins are transported until the tubules of the smooth ER eventually narrow and end. Then the proteins exit the ER in membrane-bounded, saclike organelles called **vesicles** that pinch off from the tubular endings of the membrane. Lipids are exported without a vesicle, because a vesicle is itself made of lipid.

A loaded vesicle takes its contents to the next stop in the secretory production line, the nearby **Golgi apparatus** (step 5, figure 2.5). This processing center is a stack of flat, membrane-enclosed sacs. Here, the milk sugar lactose is synthesized and other sugars are made that attach to proteins to form glycoproteins or to lipids to form glycolipids, which become parts of plasma membranes. Proteins finish folding in the Golgi apparatus.

The components of complex secretions, such as milk, are temporarily stored in the Golgi apparatus. Droplets of proteins and sugars then bud off in vesicles that move outward to the plasma membrane, fleetingly becoming part of it until they are secreted to the cell's exterior. Lipids exit the plasma membrane directly, taking bits of it with them (step 6, figure 2.5).

In the breast, epithelial cells called lactocytes form tubules, into which they secrete the components of milk. When the baby suckles, contractile cells squeeze the milk through the

tubules and out of holes in the nipples. This “ejection reflex” is so powerful that the milk can actually shoot across a room!

Intracellular Digestion— Lysosomes and Peroxisomes

Just as clutter and garbage accumulate in an apartment, debris builds up in cells. Organelles called **lysosomes** handle the garbage. Lysosomes are membrane-bounded sacs that contain enzymes that dismantle bacterial remnants, worn-out organelles, and other material such as excess cholesterol (**figure 2.6**). The enzymes also break down some digested nutrients into forms that the cell can use.

Lysosomes fuse with vesicles carrying debris from outside or within the cell, and the lysosomal enzymes then degrade the contents. For example, a type of vesicle that forms from the plasma membrane, called an endosome, ferries extra LDL cholesterol to lysosomes. A loaded lysosome moves toward the plasma membrane and fuses with it, releasing its contents to the outside. The word *lysosome* means “body that lyses;” *lyse* means “to cut.” Lysosomes maintain the very acidic environment that their enzymes require to function, without harming other cellular constituents that could be destroyed by acid.

Cells differ in number of lysosomes. Certain white blood cells and macrophages that move about and engulf bacteria are loaded with lysosomes. Liver cells require many lysosomes to break down cholesterol, toxins, and drugs.

All lysosomes contain more than 40 types of digestive enzymes, which must be maintained in a correct balance. Absence or malfunction of an enzyme causes a “lysosomal storage disease.” In these inherited disorders, which are a type of inborn error of metabolism, the molecule that the missing or abnormal enzyme normally degrades accumulates. The lysosome swells, crowding organelles and interfering with the cell's functions. In Tay-Sachs disease (MIM 272800), for example, an enzyme is deficient that normally breaks down lipids in the cells that surround nerve cells. As the nervous system becomes buried in lipid, the infant begins to lose skills, such as sight, hearing, and the ability to move. Death is typically within 3 years. Even before birth, the lysosomes of affected cells swell.

Peroxisomes are sacs with outer membranes that are studded with several types of enzymes. These enzymes perform a variety of functions, including breaking down

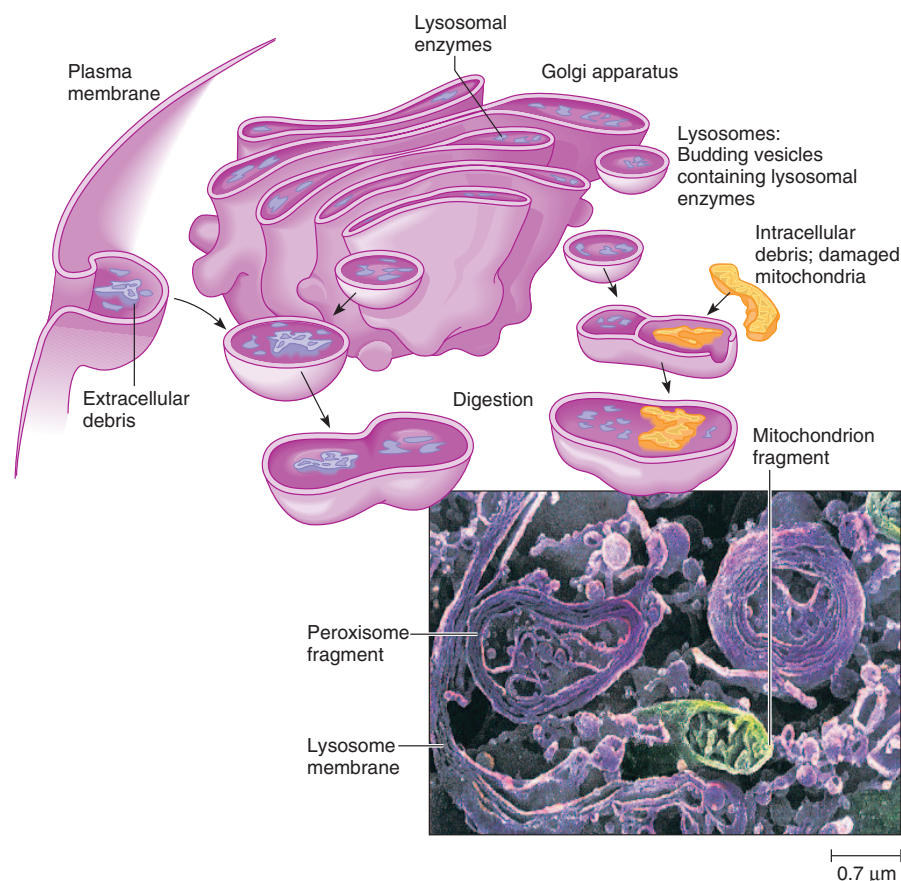


Figure 2.6 Lysosomes are trash centers. Lysosomes fuse with vesicles or damaged organelles, activating the enzymes within to recycle the molecules. Lysosomal enzymes also dismantle bacterial remnants. These enzymes require a very acidic environment to function.

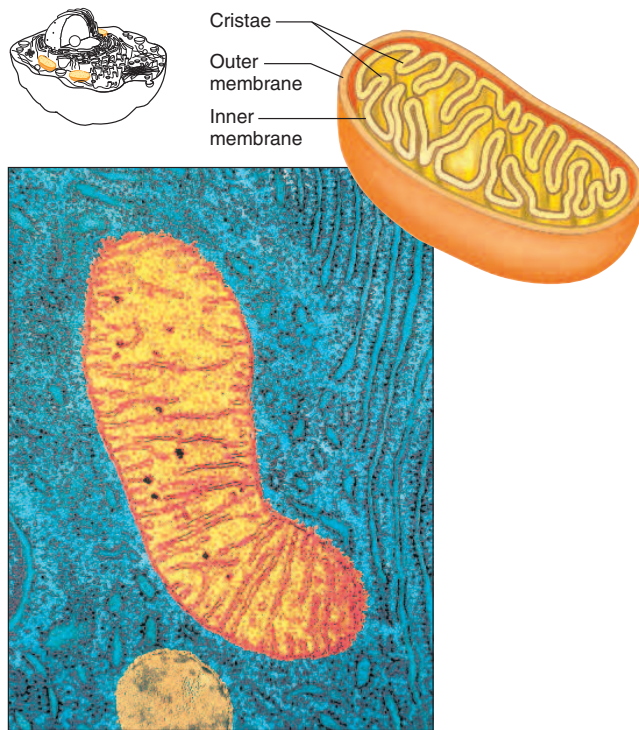


Figure 2.7 A mitochondrion extracts energy. Cristae, infoldings of the inner membrane, increase the available surface area containing enzymes for energy reactions in a mitochondrion.

certain lipids and rare biochemicals, synthesizing bile acids used in fat digestion, and detoxifying compounds that result from exposure to oxygen free radicals. Peroxisomes are large and abundant in liver and kidney cells, which handle toxins.

The 1992 film *Lorenzo's Oil* recounted the true story of a child with an inborn error of metabolism caused by an absent peroxisomal enzyme. Lorenzo had adrenoleukodystrophy (MIM202370), in which a type of lipid called a very-long-chain fatty acid builds up in the brain and spinal cord. Early symptoms include low blood sugar, skin darkening, muscle weakness, and irregular heartbeat. The patient eventually loses control over the limbs and usually dies within a few years. Eating a type of lipid in canola oil slows buildup of the very-long-chain fatty acids in blood plasma and the liver. Because the oil cannot enter the brain, eating it can only slow disease progression. A transplant of bone marrow stem cells from a compatible donor can cure the disease.

Energy Production—Mitochondria

The activities of secretion, as well as the many chemical reactions taking place in the cytoplasm, require continual energy. Organelles called **mitochondria** provide energy by breaking down nutrients from foods. The energy comes from the chemical bonds that hold together the nutrient molecules.

A mitochondrion has an outer membrane similar to those in the ER and Golgi apparatus and an inner membrane that forms folds called cristae (**figure 2.7**). These folds hold enzymes that catalyze the biochemical reactions that release energy from nutrient molecules. The energy liberated from food is captured and stored in the bonds that hold together a molecule called adenosine triphosphate (ATP). Therefore, ATP serves as a cellular energy currency.

The number of mitochondria in a cell varies from a few hundred to tens of thousands, depending upon the cell's activity level. A typical liver cell, for example, has about 1,700 mitochondria, but a muscle cell, with its very high energy requirements, has many more. Mitochondria are especially interesting because, like the nucleus, they contain DNA, although a very small amount (see figure 5.8). Chapter 5 discusses mitochondrial inheritance, and chapter 15 describes how mitochondrial genes provide insights into early human migrations.

Table 2.1 summarizes the structures and functions of organelles.

The Plasma Membrane

Just as the character of a community is molded by the people who enter and leave it, the special characteristics of different cell types are shaped in part by the substances that enter and leave. The plasma membrane controls this process. It forms a selective barrier

Table 2.1 Structures and Functions of Organelles		
Organelle	Structure	Function
Endoplasmic reticulum	Membrane network; rough ER has ribosomes, smooth ER does not	Site of protein synthesis and folding; lipid synthesis
Golgi apparatus	Stacks of membrane-enclosed sacs	Site where sugars are made and linked into starches or joined to lipids or proteins; proteins finish folding; secretions stored
Lysosome	Sac containing digestive enzymes	Degrades debris; recycles cell contents
Mitochondrion	Two membranes; inner membrane enzyme-studded	Releases energy from nutrients, participates in cell death
Nucleus	Porous sac containing DNA	Separates DNA within cell
Peroxisome	Sac containing enzymes	Breaks down and detoxifies various molecules
Ribosome	Two associated globular subunits of RNA and protein	Scaffold and catalyst for protein synthesis
Vesicle	Membrane-bounded sac	Temporarily stores or transports substances

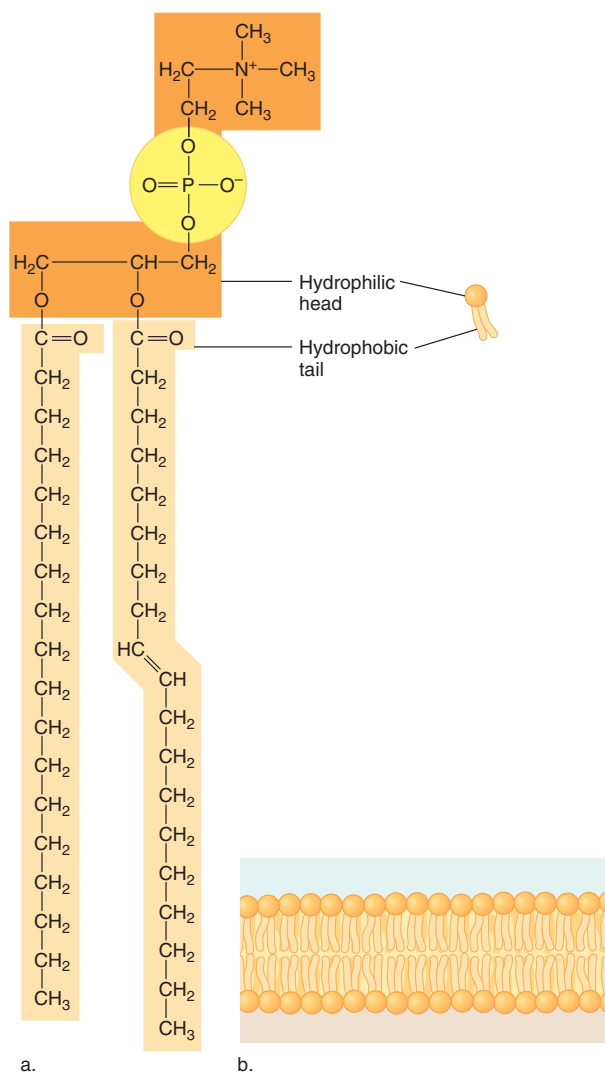


Figure 2.8 The two faces of membrane phospholipids.

(a) A phospholipid is literally a two-faced molecule, with one end attracted to water (hydrophilic, or “water-loving”) and the other repelled by it (hydrophobic, or “water-fearing”). (b) A membrane phospholipid is often depicted as a circle with two tails.

that completely surrounds the cell and monitors the movements of molecules in and out. How the chemicals that comprise the plasma membrane associate with each other determines which substances can enter or leave the cell. Membranes similar to the plasma membrane form the outer boundaries of several organelles, and some organelles consist entirely of membranes. A cell’s membranes are more than mere coverings, because some of their constituent or associated molecules carry out specific functions.

A biological membrane has a distinctive structure. It is built of a double layer (bilayer) of molecules called phospholipids. A phospholipid is a fat molecule with attached phosphate groups. It is often depicted as a head with two parallel tails. (A phosphate group $[PO_4]$ is a phosphorus atom bonded to four oxygen atoms.) Membranes can form because phospholipid

molecules self-assemble into sheets (figure 2.8). The molecules do this because their ends react oppositely to water: The phosphate end of a phospholipid is attracted to water, and thus is hydrophilic (“water-loving”); the other end, which consists of two chains of fatty acids, moves away from water, and is therefore hydrophobic (“water-fearing”). Because of these forces, phospholipid molecules in water spontaneously form bilayers. Their hydrophilic surfaces are exposed to the watery exterior and interior of the cell, and their hydrophobic surfaces face each other on the inside of the bilayer, away from water.

A phospholipid bilayer forms the structural backbone of a biological membrane. Proteins are embedded in the bilayer. Some traverse the entire structure, while others extend from a face (figure 2.9).

Proteins, glycoproteins, and glycolipids extend from a plasma membrane, creating surface topographies that are important in a cell’s interactions with other cells. The surfaces of your cells indicate not only that they are part of your body, but also that they are part of a particular organ and a particular tissue type.

Many molecules that extend from the plasma membrane are **receptors**, which are structures that have indentations or other shapes that fit and hold molecules outside the cell. The molecule that binds to the receptor, called the **ligand**, may set into motion a cascade of chemical reactions that carries out a particular cellular activity, such as dividing.

The phospholipid bilayer is oily, and some proteins move within it like ships on a sea. Proteins with related functions may cluster on “lipid rafts” that float on the phospholipid bilayer. The rafts are rich in cholesterol and other types of lipids. This clustering of proteins eases their interaction.

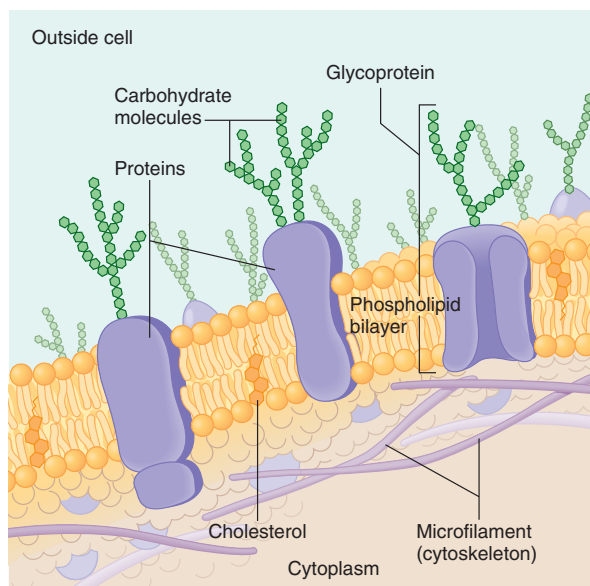


Figure 2.9 Anatomy of a plasma membrane. In a plasma membrane, mobile proteins are embedded throughout a phospholipid bilayer. Other types of lipids aggregate to form “rafts,” and an underlying mesh of protein fibers provides support. Carbohydrates jut from the membrane’s outer face.

Proteins aboard lipid rafts have several functions. They contribute to the cell's identity; act as transport shuttles into the cell; serve as gatekeepers; and can let in certain toxins and pathogens. HIV, for example, enters a cell by breaking a lipid raft.

The inner hydrophobic region of the phospholipid bilayer blocks entry and exit to most substances that dissolve in water.

However, certain molecules can cross the membrane through proteins that form passageways, or when they are escorted by a "carrier" protein. Some membrane proteins form channels for ions, which are atoms or molecules with an electrical charge. **Reading 2.2** describes "channelopathies"—diseases that stem from faulty ion channels.

Reading 2.2

Faulty Ion Channels Cause Inherited Disease

What do abnormal pain intensity, irregular heartbeats, and cystic fibrosis have in common? All result from abnormal ion channels in plasma membranes.

Ion channels are protein-lined tunnels in the phospholipid bilayer of a biological membrane. These passageways permit electrical signals in the form of ions (charged particles) to pass through membranes.

Ion channels are specific for calcium (Ca^{+2}), sodium (Na^{+}), potassium (K^{+}), or chloride (Cl^{-}) ions. A plasma membrane may have a few thousand ion channels for each of these ions. Ten million ions can pass through an ion channel in one second! The following "channelopathies" result from abnormal ion channels.

Absent or Extreme Pain

The 10-year-old boy amazed the people on the streets of his small, northern Pakistani town. He was completely unable to feel pain, so he had become a performer, stabbing knives through his arms and walking on hot coals to entertain crowds. Several other people in this community where relatives often married relatives were also unable to feel pain. Researchers studied the connected families and discovered a mutation that alters sodium channels on certain nerve cells. The mutation blocks the channels so that the message to feel pain cannot be sent. The boy died at age 13 from jumping off a roof. His genes could protect him from pain, but pain protects against injury by providing a warning. He foolishly jumped.

A different mutation affecting the same sodium channel causes very different symptoms. In "burning man syndrome," the channels become hypersensitive, opening and flooding the body with pain easily, in response to exercise, an increase in room temperature, or just putting on socks. In another condition, "paroxysmal extreme pain disorder," the sodium channels stay open too long, causing excruciating pain in the rectum, jaw, and eyes. Researchers are using the information from studies of these genetic disorders to develop new painkillers.

Long-QT Syndrome and Potassium Channels

Four children in a Norwegian family were born deaf, and three of them died at ages 4, 5, and 9. All of the children had inherited from unaffected carrier parents "long-QT syndrome associated with deafness" (MIM 176261). ("QT" refers to part of a normal heart rhythm.) These children had abnormal potassium channels in the cells of the heart muscle and in the inner ear. In the heart cells, the malfunctioning ion channels

disrupted electrical activity, fatally disturbing heart rhythm. In the cells of the inner ear, the abnormal ion channels increased the extracellular concentration of potassium ions, impairing hearing.

Cystic Fibrosis and Chloride Channels

A seventeenth-century English saying, "A child that is salty to taste will die shortly after birth," described the consequence of abnormal chloride channels in CF. The chloride channel is called CFTR, for cystic fibrosis transductance regulator. In most cases, CFTR protein remains in the cytoplasm, unable to reach the plasma membrane, where it would normally function (**figure 1**).

CF is inherited from carrier parents. The major symptoms of difficulty breathing, frequent severe respiratory infections, and a clogged pancreas that disrupts digestion all result from a buildup of extremely thick mucous secretions.

Abnormal chloride channels in cells lining the lung passageways and ducts of the pancreas cause the symptoms of CF. The primary defect in the chloride channels also disrupts sodium channels. The result: Salt trapped inside cells draws moisture in and thickens surrounding mucus.

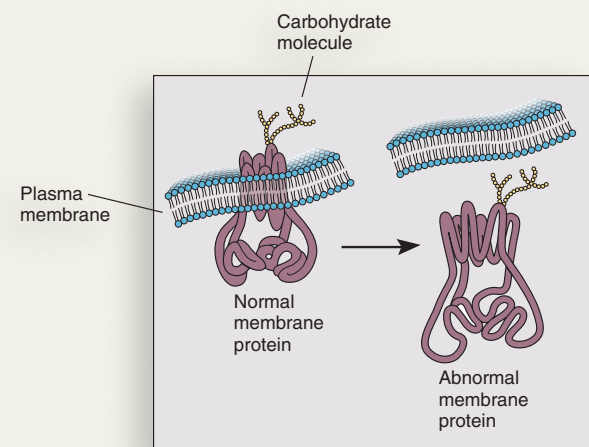


Figure 1 In cystic fibrosis, CFTR protein remains in the cytoplasm, rather than anchoring in the plasma membrane. This prevents normal chloride channel function.

The Cytoskeleton

The **cytoskeleton** is a meshwork of protein rods and tubules that molds the distinctive structures of a cell, positioning organelles and providing three-dimensional shape. The proteins of the cytoskeleton are continually broken down and built up as a cell performs specific activities. Some cytoskeletal elements function as rails, forming conduits that transport cellular contents; other parts, called motor molecules, power the movement of organelles along these rails by converting chemical energy to mechanical energy.

The cytoskeleton includes three major types of elements—**microtubules**, **microfilaments**, and **intermediate filaments** (figure 2.10). They are distinguished by protein type, diameter, and how they aggregate into larger structures. Other proteins connect these components, creating the framework that provides the cell's strength and ability to resist force and maintain shape.

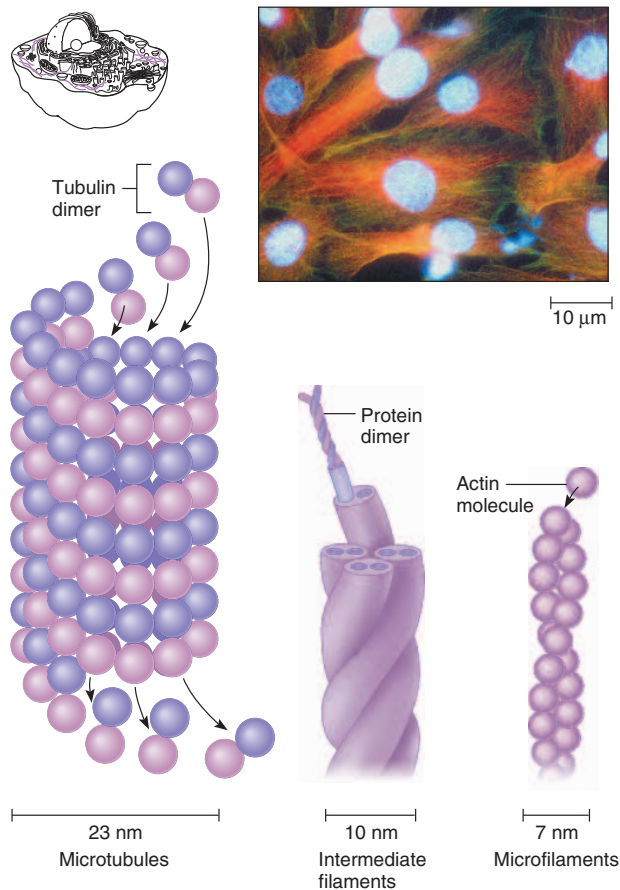


Figure 2.10 The cytoskeleton is made of protein rods and tubules. The three major components of the cytoskeleton are microtubules, intermediate filaments, and microfilaments. Through special staining, the cytoskeletons in these cells appear orange under the microscope. (The abbreviation nm stands for nanometer, which is a billionth of a meter.)

Long, hollow microtubules provide many cellular movements. A microtubule is composed of pairs (dimers) of a protein, called tubulin, assembled into a hollow tube. The cell can change the length of the tubule by adding or removing tubulin molecules.

Cells contain both formed microtubules and individual tubulin molecules. When the cell requires microtubules to carry out a specific function—cell division, for example—free tubulin dimers self-assemble into more tubules. After the cell divides, some of the microtubules fall apart into individual tubulin dimers, replenishing the cell's supply of building blocks. Cells are perpetually building up and breaking down microtubules. Some drugs used to treat cancer affect the microtubules that pull a cell's duplicated chromosomes apart, either by preventing tubulin from assembling into microtubules, or by preventing microtubules from breaking down into free tubulin dimers. In each case, cell division stops.

Microtubules also form cilia, which are hairlike structures (figure 2.11). Coordinated movement of cilia generates a wave that moves the cell or propels substances along its surface. Cilia beat particles up and out of respiratory tubules, and cilia move egg cells in the female reproductive tract. Because cilia are so widespread in the body, defects in them affect health. One such “ciliopathy” is Bardet-Biedl syndrome (MIM 209900), which causes obesity, visual loss, diabetes, cognitive impairment, and extra fingers and/or toes.

Microfilaments, are long, thin rods composed of many molecules of the protein actin. They are solid and narrower than microtubules, enable cells to withstand stretching and compression, and help anchor one cell to another. Microfilaments provide many other functions in the cell through proteins that interact with actin. When any of these proteins is absent or abnormal, a genetic disease results.

Intermediate filaments have diameters intermediate between those of microtubules and microfilaments, and are made of different proteins in different cell types. However, all intermediate filaments share a common overall organization of dimers entwined into nested coiled rods. Intermediate filaments are scarce in many cell types but are very abundant in skin and nerve cells.

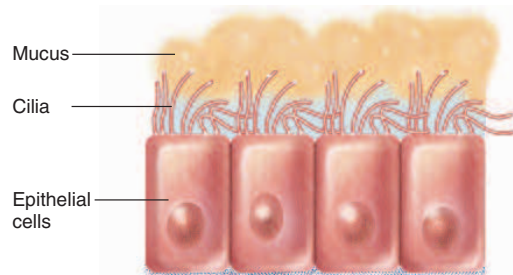


Figure 2.11 Microtubules form cilia, which are hairlike structures that wave, moving secretions such as mucus on the cell surfaces.

The intermediate filaments in actively dividing skin cells in the bottommost layer of the epidermis (the upper skin layer) form a strong inner framework that firmly attaches the cells to each other and to the underlying tissue. These cellular attachments are crucial to the skin's barrier function. In a group of inherited conditions called epidermolysis bullosa (MIM 226500, 226650, 131750), intermediate filaments are abnormal. The skin blisters easily as tissue layers separate. The “*In Their Own Words*” essay describes how abnormal intermediate filaments affect a little girl, who has giant axonal neuropathy (MIM 256850).

Disruption of how the cytoskeleton interacts with other cell components can be devastating. Consider hereditary spherocytosis (MIM 182900), which disturbs the interface

between the plasma membrane and the cytoskeleton in red blood cells.

The doughnut shape of normal red blood cells enables them to squeeze through the narrowest blood vessels. Their cytoskeletons provide the ability to deform. Rods of a protein called spectrin form a meshwork beneath the plasma membrane, strengthening the cell. Proteins called ankyrins attach the spectrin rods to the plasma membrane (**figure 2.12**). Spectrin molecules also attach to microfilaments and microtubules. Spectrin molecules are like steel girders, and ankyrins are like nuts and bolts. If either molecule is absent, the red blood cell collapses.

In hereditary spherocytosis, the ankyrins are abnormal, and parts of the red blood cell plasma membrane disintegrate.

In Their Own Words

A Little Girl with Giant Axons

A nerve cell (neuron) communicates by receiving electrochemical signals at one highly branched end, and sending signals from the other end, which is a single branch called an axon. Intermediate filaments, called neurofilaments, control the axon's shape. In giant axonal neuropathy (GAN), a key neurofilament protein, gigaxonin, is not dismantled and recycled as it normally is, and instead builds up in axons, distending them. The giant axons stifle nerve transmission, affecting the ability to move, sense, and think. A little-understood but striking part of the phenotype is very curly hair. An affected individual is wheelchair-bound by adolescence, and does not survive his or her twenties. Lori Sames tells about her daughter, Hannah, who has GAN.

“Hannah Sarah Sames is a beautiful little girl who was born on March 5, 2004. She has extremely curly blonde hair, a slight build, a precocious smile, and a charming personality. She loves to sing and dance, and play outdoors. Hannah is a beaming light of love.

When Hannah was 2 years, 5 months old, her grandmother noticed her left arch seemed to be rolling inward. I took Hannah to an orthopedist and a podiatrist, and was told Hannah would be fine. But by her third birthday, we suspected something was wrong—both arches were now involved, and her gait had become awkward. Her pediatrician gave her a rigorous physical exam and agreed she had an awkward gait, but felt that was just how Hannah walks.

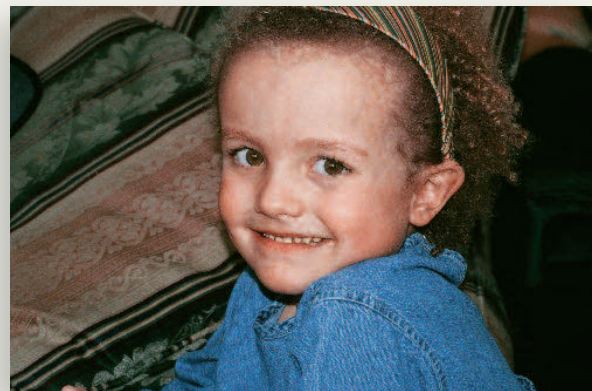
Two months later, I took Hannah to another orthopedist, who told me to just let her live her life, she would be fine. Convinced otherwise, my sister showed cell phone video of Hannah walking to a physical therapist she works with, who thought Hannah's gait was like that of a child with muscular dystrophy. Our pediatrician referred us to a pediatric neurologist and a pediatric geneticist, and 6 months of testing for various diseases began. Results: all normal. During another visit with the pediatric neurologist, he took out a huge textbook and showed us a photo of a skinny little boy with kinky hair and a high forehead and braces that went just below the knee—he had GAN. He looked exactly like Hannah. So off we went to a children's hospital in New York City for more tests, and the diagnosis of GAN was confirmed.

Meeting with a genetic counselor 3 days later brought devastation. Matt and I are each carriers, and we passed the disease to Hannah. Each of our two other daughters has a 2 in 3 chance of being a carrier. We learned GAN is a rare “orphan genetic disorder” for which there is no cure, no treatment, no clinical trial and no ongoing research. ‘So you are telling us this is a death sentence?’ I asked. And, we were told, ‘Yes’.

Matt and I walked around in a state of shock, anger, disbelief, and grief for two days. Then, we realized, as with any disease, someone has to be the first to be cured. Some family has to be the first to raise funds and awareness and pull the medical community together to find treatment. This is how Hannah's Hope Foundation was born! As a result, we held the world's first symposium for GAN, where clinicians and scientists brainstormed. Our foundation is now funding a number of projects aimed at treating GAN.”

Lori Sames

<http://www.hannahshopefund.org/>



Hannah Sames has giant axonal neuropathy, a disorder that affects intermediate filaments in nerve cells. Her beautiful curls are one of the symptoms.

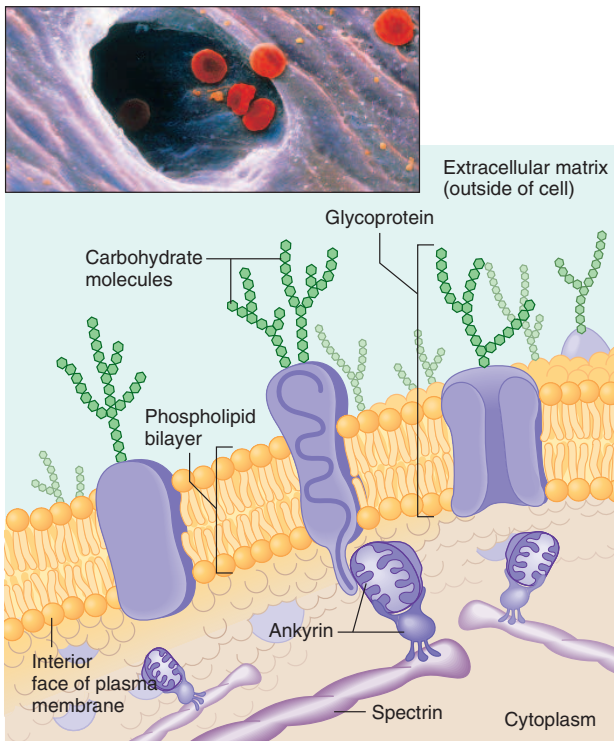


Figure 2.12 The red blood cell plasma membrane. The cytoskeleton that supports the plasma membrane of a red blood cell withstands the turbulence of circulation. Proteins called ankyrins bind molecules of spectrin from the cytoskeleton to the inner membrane surface. On its other end, ankyrin binds proteins that help ferry molecules across the plasma membrane. In hereditary spherocytosis, abnormal ankyrin collapses the plasma membrane. The cell balloons—a problem for a cell whose function depends upon its shape. The inset shows normal red blood cells.

The cell balloons out, obstructing narrow blood vessels—especially in the spleen, the organ that normally disposes of aged red blood cells. Anemia develops as the spleen destroys red blood cells more rapidly than the bone marrow can replace them. The result is great fatigue and weakness. Removing the spleen can treat the condition.

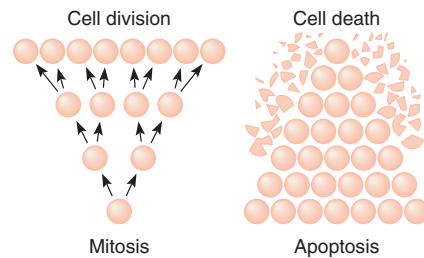
Key Concepts

1. Cells are the units of life. They consist mostly of carbohydrates, lipids, proteins, and nucleic acids.
2. Organelles subdivide specific cell functions. They include the nucleus, the endoplasmic reticulum (ER), Golgi apparatus, mitochondria, lysosomes, and peroxisomes.
3. The plasma membrane is a flexible, selective phospholipid bilayer with embedded proteins and lipid rafts.
4. The cytoskeleton is an inner framework made of protein rods and tubules, connectors and motor molecules.

2.3 Cell Division and Death

A human body is not a static object with a set number of cells. Instead, new cells are continually forming, and old ones dying, at different rates in different tissues. Growth, development, maintaining health, and healing from disease or injury require an intricate interplay between the rates of these two processes: **mitosis**, a form of cell division that gives rise to two somatic cells from one, and **apoptosis**, a form of cell death (figure 2.13).

About 10 trillion of a human body's 100 or so trillion cells are replaced daily. Yet, cell death must happen to mold certain organs, just as a sculptor must remove some clay to shape the desired object. Apoptosis carves toes, for example, from weblike structures that telescope out from an embryo's developing form. Apoptosis, which comes from the Greek for “leaves falling from a tree,” is a precise, genetically programmed sequence of events that is a normal part of development.



a.



b.

Figure 2.13 Mitosis and apoptosis mold a body. (a) Cell numbers increase from mitosis and decrease from apoptosis. (b) In the embryo, fingers and toes are carved from webbed structures. In syndactyly, normal apoptosis fails to carve digits, and webbing persists.

The Cell Cycle

Many cell divisions transform a fertilized egg into a many-trillion-celled person. A series of events called the **cell cycle** describes the sequence of activities as a cell prepares for division and then divides.

Cell cycle rate varies in different tissues at different times. A cell lining the small intestine's inner wall may divide throughout life, whereas a neuron in the brain may never divide; a cell in the deepest skin layer of a 90-year-old may divide as long as the person lives. Frequent mitosis enables the embryo and fetus to grow rapidly. By birth, the mitotic rate slows dramatically. Later, mitosis maintains the numbers and positions of specialized cells in tissues and organs.

The cell cycle is continual, but we divide it into stages based on what we observe. The two major stages are **interphase** (not dividing) and **mitosis** (dividing) (**figure 2.14**). In mitosis, a cell duplicates its chromosomes, then apportions one set into each of two resulting cells, called daughter cells. This maintains the set of 23 chromosome pairs characteristic of a human somatic cell. Another form of cell division, meiosis, produces sperm or eggs, which have half the amount of genetic material in somatic cells, or 23 single chromosomes. Chapter 3 discusses meiosis.

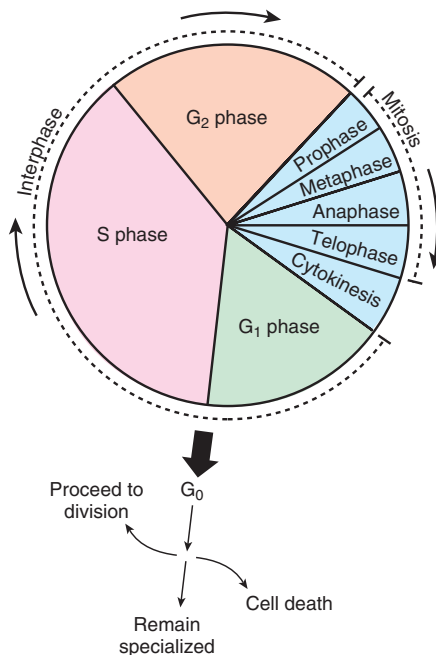


Figure 2.14 The cell cycle. The cell cycle is divided into interphase, when cellular components are replicated, and mitosis, when the cell distributes its contents into two daughter cells. Interphase is divided into G₁ and G₂, when the cell duplicates specific molecules and structures, and S phase, when it replicates DNA. Mitosis is divided into four stages plus cytokinesis, when the cells separate. G₀ is a “time-out” when a cell “decides” which course of action to follow.

Interphase—A Time of Great Activity

Interphase is a very active time. The cell continues the basic biochemical functions of life and also replicates its DNA and other subcellular structures. Interphase is divided into two gap (**G₁** and **G₂**) **phases** and one synthesis (**S**) **phase**. In addition, a cell can exit the cell cycle at G₁ to enter a quiescent phase called **G₀**. A cell in G₀ maintains its specialized characteristics but does not replicate its DNA or divide. From G₀, a cell may also proceed to mitosis and divide, or die. Apoptosis may ensue if the cell's DNA is so damaged that cancer might result. G₀, then, is when a cell's fate is either decided or put on hold.

During G₁, which follows mitosis, the cell resumes synthesis of proteins, lipids, and carbohydrates. These molecules will contribute to building the extra plasma membrane required to surround the two new cells that form from the original one. G₁ is the period of the cell cycle that varies the most in duration among different cell types. Slowly dividing cells, such as those in the liver, may exit at G₁ and enter G₀, where they remain for years. In contrast, the rapidly dividing cells in bone marrow speed through G₁ in 16 to 24 hours. Cells of the early embryo may skip G₁ entirely.

During S phase, the cell replicates its entire genome. As a result, each chromosome then consists of two copies joined at an area called the **centromere**. In most human cells, S phase takes 8 to 10 hours. Many proteins are also synthesized during this phase, including those that form the mitotic **spindle** that will pull the chromosomes apart. Microtubules form structures called **centrioles** near the nucleus. Centriole microtubules join with other proteins and are oriented at right angles to each other, forming paired, oblong structures called **centrosomes** that organize other microtubules into the spindle.

Mutations in genes that encode proteins of the centrosome cause microcephaly, in which the brain is very small but intelligence may be normal. The connection between impaired cell division and a small brain is not known.

G₂ occurs after the DNA has been replicated but before mitosis begins. More proteins are synthesized during this phase. Membranes are assembled from molecules made during G₁ and are stored as small, empty vesicles beneath the plasma membrane. These vesicles will merge with the plasma membrane to enclose the two daughter cells.

Mitosis—The Cell Divides

As mitosis begins, the replicated chromosomes are condensed enough to be visible, when stained, under a microscope. The two long strands of identical chromosomal material in a replicated chromosome are called **chromatids** (**figure 2.15**). At a certain point during mitosis, a replicated chromosome's centromere splits, allowing its chromatid pair to separate into two individual chromosomes. (Although the centromere of a replicated chromosome appears as a constriction, its DNA is replicated.)

During **prophase**, the first stage of mitosis, DNA coils tightly. This shortens and thickens the chromosomes, which enables them to more easily separate (**figure 2.16**). Microtubules assemble from tubulin building blocks in the cytoplasm

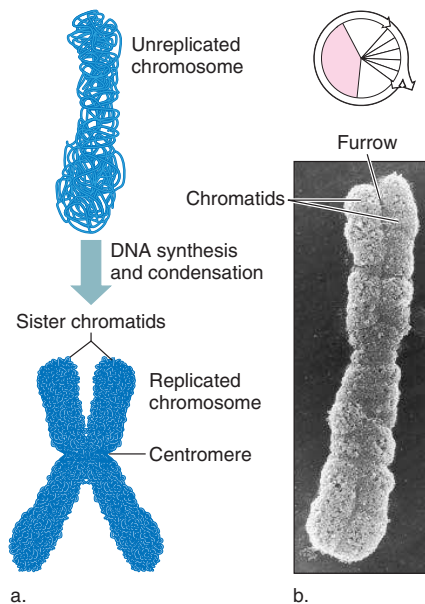


Figure 2.15 Replicated and unreplicated chromosomes. Chromosomes are replicated during S phase, before mitosis begins. Two genetically identical chromatids of a replicated chromosome join at the centromere (a). In the photograph (b), a human chromosome is forming two chromatids.

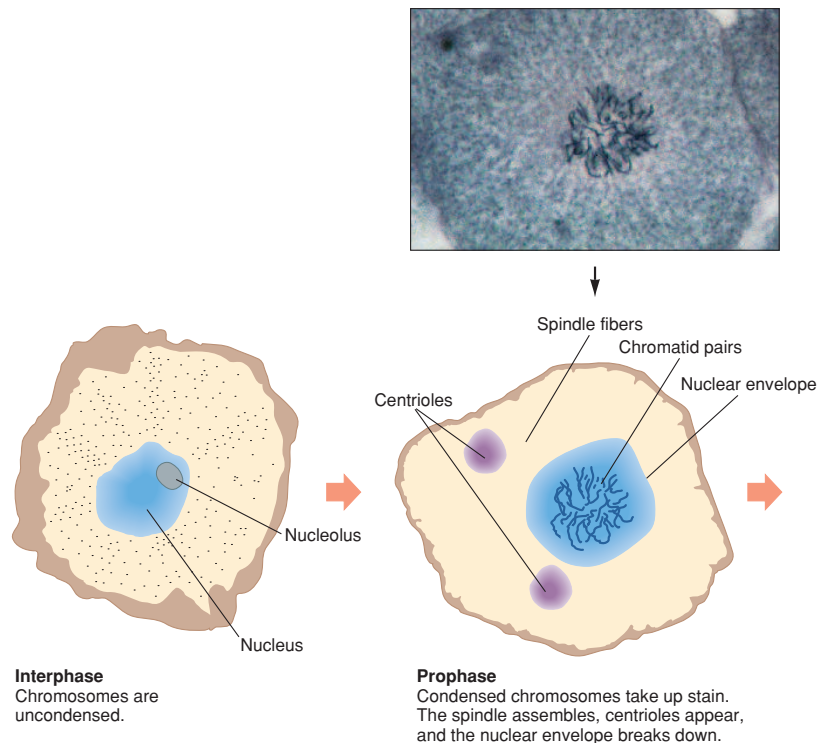


Figure 2.16 Mitosis in a human cell. Replicated chromosomes separate and are distributed into two cells from one. In a separate process, cytokinesis, the cytoplasm and other cellular structures distribute and pinch off into two daughter cells. (Not all chromosome pairs are depicted.)

to form the spindles. Toward the end of prophase, the nuclear membrane breaks down. The nucleolus is no longer visible.

Metaphase follows prophase. Chromosomes attach to the spindle at their centromeres and align along the center of the cell, which is called the equator. Metaphase chromosomes are under great tension, but they appear motionless because they are pulled with equal force on both sides, like a tug-of-war rope pulled taut.

Next, during **anaphase**, the plasma membrane indents at the center, where the metaphase chromosomes line up. A band of microfilaments forms on the inside face of the plasma membrane, constricting the cell down the middle. Then the centromeres part, which relieves the tension and releases one chromatid from each pair to move to opposite ends of the cell—like a tug-of-war rope breaking in the middle and the participants falling into two groups. Microtubule movements stretch the dividing cell. During the very brief anaphase stage, a cell temporarily contains twice the normal number of chromosomes because each chromatid becomes an independently moving chromosome, but the cell has not yet physically divided.

In **telophase**, the final stage of mitosis, the cell looks like a dumbbell with a set of chromosomes at each end. The spindle falls apart, and nucleoli and the membranes around the nuclei re-form at each end of the elongated cell. Division of the genetic material is now complete. Next, during **cytokinesis**, organelles and macromolecules are distributed between the two daughter

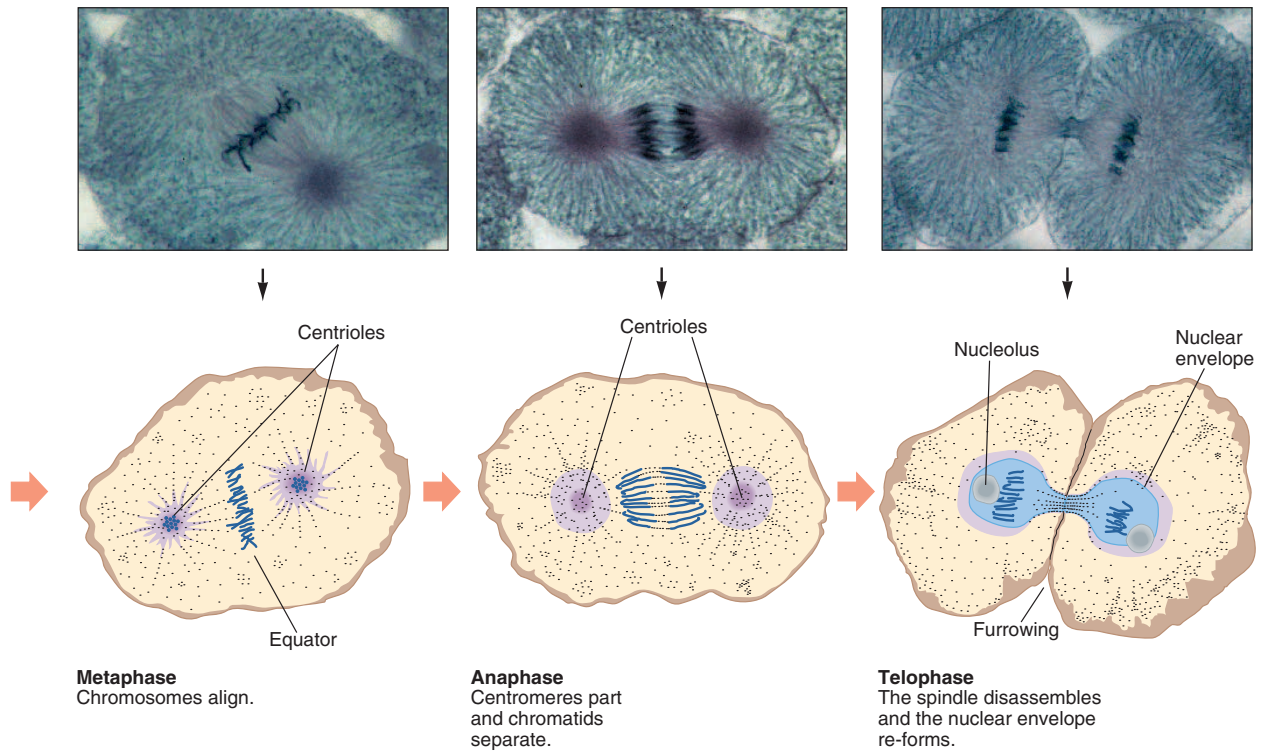
cells. Finally, the microfilament band contracts like a drawstring, separating the newly formed cells.

Control of the Cell Cycle

When and where a somatic cell divides is crucial to health. Illness can result from abnormally regulated mitosis. Control of mitosis is a daunting task. Quadrillions of mitoses occur in a lifetime, and not at random. Too little mitosis, and an injury goes unrepaired; too much, and an abnormal growth forms.

Groups of interacting proteins function at times in the cell cycle called checkpoints to ensure that chromosomes are faithfully replicated and apportioned into daughter cells (figure 2.17). A “DNA damage checkpoint,” for example, temporarily pauses the cell cycle while special proteins repair damaged DNA. An “apoptosis checkpoint” turns on as mitosis begins. During this checkpoint, proteins called survivins override signals telling the cell to die, ensuring that mitosis (division) rather than apoptosis (death) occurs. Later during mitosis, the “spindle assembly checkpoint” oversees construction of the spindle and the binding of chromosomes to it.

Cells obey an internal “clock” that tells them approximately how many times to divide. Mammalian cells grown (cultured) in a dish divide about 40 to 60 times. The mitotic clock ticks down with time. A connective tissue cell from a



fetus, for example, will divide about 50 more times. A similar cell from an adult divides only 14 to 29 more times.

How can a cell “know” how many divisions remain? The answer lies in the chromosome tips, called **telomeres** (figure 2.18). Telomeres function like cellular fuses that burn down as pieces are lost from the ends. Telomeres consist of hundreds to thousands of repeats of a specific six DNA-base

sequence. At each mitosis, the telomeres lose 50 to 200 end-most bases, gradually shortening the chromosome. After about 50 divisions, a critical length of telomere DNA is lost, which signals mitosis to stop. The cell may remain alive but not divide again, or it may die.

Not all cells have shortening telomeres. In eggs and sperm, in cancer cells, and in a few types of normal cells that must continually supply new cells (such as bone marrow cells),

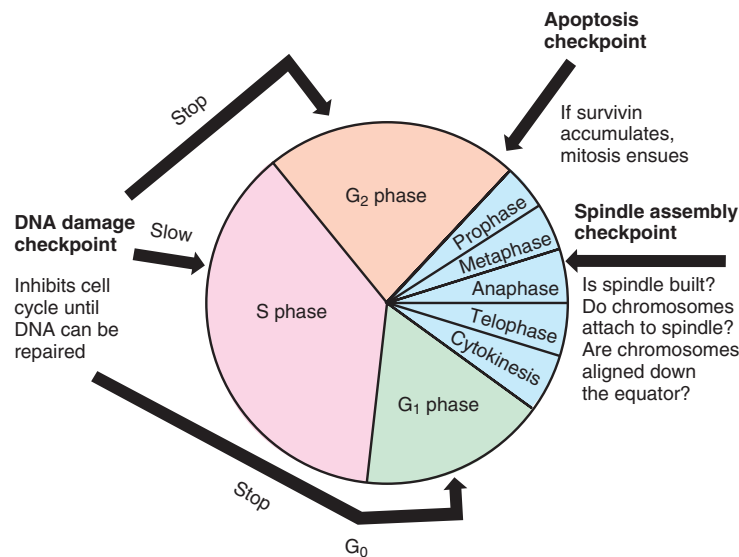


Figure 2.17 Cell cycle checkpoints. Checkpoints ensure that mitotic events occur in the correct sequence. Many types of cancer result from faulty checkpoints.

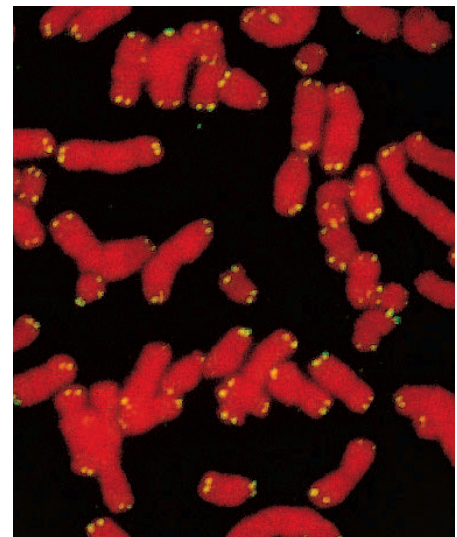


Figure 2.18 Telomeres. Fluorescent tags mark the telomeres in this human cell.

an enzyme called telomerase keeps chromosome tips long (see figure 18.4). However, most cells do not produce telomerase, and their chromosomes gradually shrink.

The rate of telomere shortening provides a “clock” for a cell’s existence. The telomere clock may not only count down a cell’s remaining lifespan, but may also sense environmental stimuli. For example, chronic stress, obesity, and elevated blood sugar are associated with accelerated telomere shortening. It may be possible to alter the environment to keep telomeres longer. One study compared healthy identical twins, one of whom was physically active and the other sedentary. The twins who exercised had on average 200 more DNA bases on their chromosome tips than did the sedentary twins.

Factors from outside the cell can also affect a cell’s mitotic clock. Crowding can slow or halt mitosis. Normal cells growing in culture stop dividing when they form a one-cell-thick layer lining the container. This limitation to division is called contact inhibition. If the layer tears, the cells that border the tear grow and divide, filling in the gap. They stop dividing once the space is filled. Perhaps a similar mechanism in the body limits mitosis.

Chemical signals control the cell cycle from outside. Hormones and growth factors are biochemicals from outside the cell that influence mitotic rate. A **hormone** is a substance synthesized in a gland and transported in the bloodstream to another part of the body, where it exerts a specific effect. Hormones secreted in the brain, for example, signal the cells lining a woman’s uterus to build up each month by mitosis in preparation for possible pregnancy. A growth factor acts more locally. Epidermal growth factor (EGF), for example, stimulates cell division in the skin beneath a scab. A class of cancer drugs works by plugging up EGF receptors on cancer cells, blocking incoming signals to divide.

Two types of proteins, cyclins and kinases, interact inside cells to activate the genes whose products carry out mitosis. The two types of proteins form pairs. Cyclin levels fluctuate regularly throughout the cell cycle, while kinase levels stay the same. A certain number of cyclin-kinase pairs turn on the genes that trigger mitosis. Then, as mitosis begins, enzymes degrade the cyclin. The cycle starts again as cyclin begins to build up during the next interphase.

Apoptosis

Apoptosis rapidly and neatly dismantles a cell into membrane-enclosed pieces that a phagocyte (a cell that engulfs and destroys another) can mop up. It is a little like packing the contents of a messy room into garbage bags, then disposing of it all. In contrast is necrosis, a form of cell death associated with inflammation, rather than an orderly, contained destruction.

Like mitosis, apoptosis is continuous. It begins when a “death receptor” on the cell’s plasma membrane receives a signal

to die. Within seconds, enzymes called caspases are activated inside the doomed cell, stimulating each other and snipping apart various cell components. These killer enzymes:

- Demolish enzymes that replicate and repair DNA.
- Activate enzymes that cut DNA into similarly sized small pieces.
- Tear apart the cytoskeleton, including threads that support the nucleus, which collapses, condensing the DNA within.
- Cause mitochondria to release molecules that trigger further caspase activity, end the cell’s energy supply, and destroy these organelles.
- Abolish the cell’s ability to adhere to other cells.
- Send a certain phospholipid from the plasma membrane’s inner face to its outer surface, where it attracts phagocytes that dismantle the cell remnants.

A dying cell has a characteristic appearance (figure 2.19). It rounds up as contacts with other cells are cut off, and the plasma membrane undulates, forming bulges called

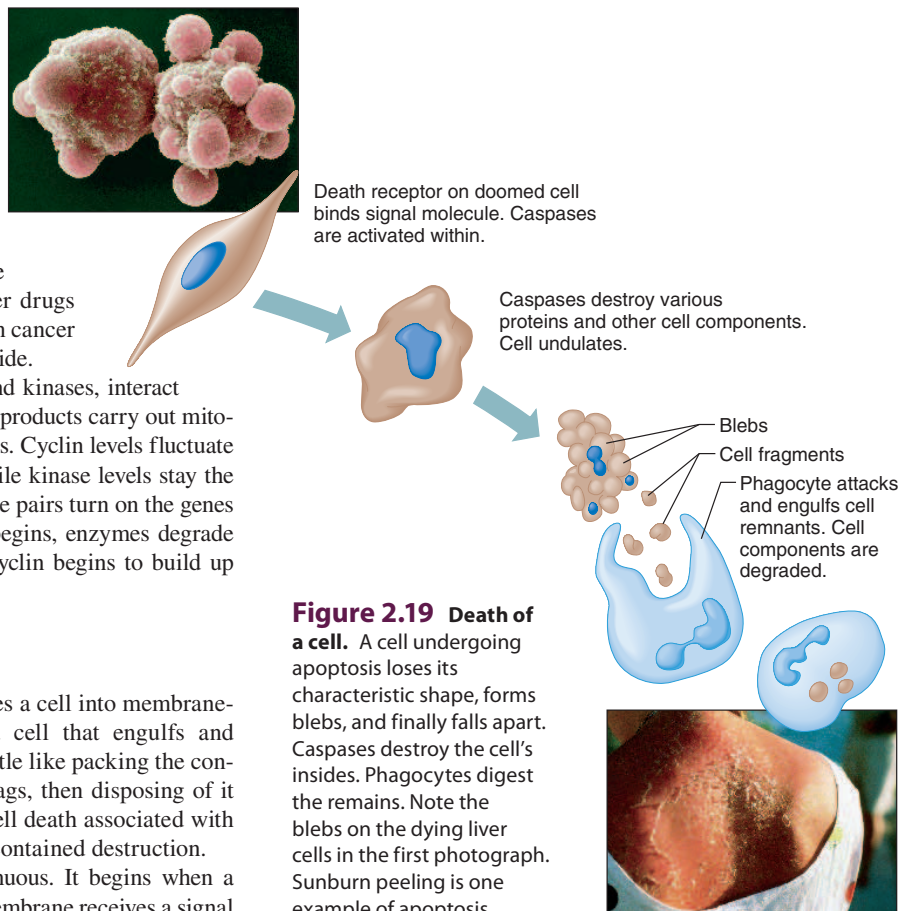


Figure 2.19 **Death of a cell.** A cell undergoing apoptosis loses its characteristic shape, forms blebs, and finally falls apart. Caspases destroy the cell’s insides. Phagocytes digest the remains. Note the blebs on the dying liver cells in the first photograph. Sunburn peeling is one example of apoptosis.

blebs. The nucleus bursts, releasing same-sized DNA pieces. Mitochondria decompose. Then the cell shatters. Almost instantly, pieces of membrane encapsulate the cell fragments, which prevents inflammation. Within an hour, the cell is gone.

From the embryo onward through development, mitosis and apoptosis are synchronized, so that tissue neither overgrows nor shrinks. In this way, a child's liver retains much the same shape as she grows into adulthood, yet enlarges. During early development, mitosis and apoptosis orchestrate the ebb and flow of cell number as new structures form. Later, these processes protect—mitosis produces new skin to heal a scraped knee; apoptosis peels away sunburnt skin cells that might otherwise become cancerous. Cancer is a profound derangement of the balance between cell division and death. In cancer, mitosis occurs too frequently or too many times, or apoptosis happens too infrequently. Chapter 18 discusses cancer in detail.

Key Concepts

1. Mitosis and apoptosis regulate cell numbers during development, growth, and repair.
2. The cell cycle includes interphase and mitosis. During G_0 , the cell "decides" to divide, die, or stay differentiated. Interphase includes two gap (G) phases and a synthesis (S) phase that prepares the cell for mitosis. During S phase, DNA is replicated. Proteins, carbohydrates, and lipids are synthesized during G_1 and more proteins are synthesized in G_2 . During mitosis, replicated chromosomes condense, align, split, and distribute into daughter cells.
3. The cell cycle is controlled by checkpoints; telomeres; hormones and growth factors from outside the cell; and cyclins and kinases from within.
4. During apoptosis, cells receive a death signal, activate caspases, and break apart in an orderly fashion.

2.4 Cell-Cell Interactions

Precisely coordinated biochemical steps orchestrate the cell-cell interactions that make multicellular life possible. Defects in cell communication and interaction cause certain inherited illnesses. Two broad types of interactions among cells are signal transduction and cellular adhesion.

Signal Transduction

In **signal transduction**, molecules on the plasma membrane assess, transmit, and amplify incoming messages to the cell's interior. *Transduce* means to change one form of something (such as energy or information) into another. In signal transduction, the cell changes various types of stimuli into specific biochemical reactions. A cell's existence may depend upon particular signal molecules binding receptors on the cell surface. Yet other signals must be ignored for cell

survival, such as a signal to divide when cell division is not warranted. A cell's response to the many signals it receives is very complex.

The proteins that carry out signal transduction are in the cytoplasm and are embedded in the plasma membrane, from which they extend from one or both faces. The proteins act in a sequence, beginning at the cell surface. First, a receptor binds an incoming molecule, called the "first messenger." The receptor then contorts, touching a nearby protein called a regulator (figure 2.20). Next, the regulator activates a nearby enzyme, which catalyzes (speeds) a specific chemical reaction. The product of this reaction, called the "second messenger," is the key part of the entire process because it elicits the cell's response. This is usually activation of certain enzymes.

A single stimulus can trigger the production of many second messenger molecules. This is how signal transduction amplifies incoming information. Because cascades of proteins carry out signal transduction, it is a genetically controlled process.

Defects in signal transduction underlie many inherited disorders. In neurofibromatosis type 1 (NF1) (MIM 162200), for example, tumors (usually benign) grow in nervous tissue, particularly under the skin. At the cellular level, NF1 occurs when cells fail to block transmission of a growth factor signal that triggers cell division. Affected cells misinterpret the signal and divide when it is inappropriate.

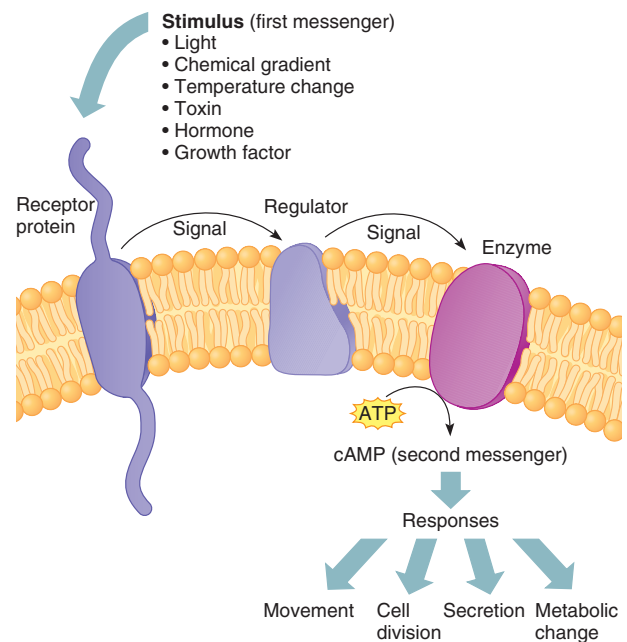


Figure 2.20 Signal transduction. A receptor binds a first messenger, triggering a cascade of biochemical activity at the cell's surface. An enzyme catalyzes a reaction inside the cell that circularizes ATP to cyclic AMP, the second messenger. cAMP then stimulates various responses, such as cell division, metabolic changes, and muscle contraction. Splitting ATP also releases energy.

Cellular Adhesion

Cellular adhesion is a precise sequence of interactions among the proteins that connect cells. Inflammation—the painful, red swelling at a site of injury or infection—illustrates one type of cellular adhesion. Inflammation occurs when white blood cells (leukocytes) move in the circulation to the injured or infected body part. There they squeeze between cells of the blood vessel walls to exit the circulation and reach the injury site. Cellular adhesion molecules, or CAMs, help guide white blood cells to the injured area.

Three types of CAMs carry out the inflammatory response: selectins, integrins, and adhesion receptor proteins (**figure 2.21**). First, selectins attach to the white blood cells and slow them to a roll by also binding to carbohydrates on the capillary wall. (This is a little like putting out your arms to slow your ride down a slide.) Next, clotting blood, bacteria, or decaying tissues release chemical attractants that signal white blood cells to stop. The chemical attractants activate CAMs called integrins, which latch onto the white blood cells, and CAMs called adhesion receptor proteins, which extend from the capillary wall at the injury site. The integrins and adhesion receptor proteins then guide the white blood cells between the tile-like lining cells to the injury site.

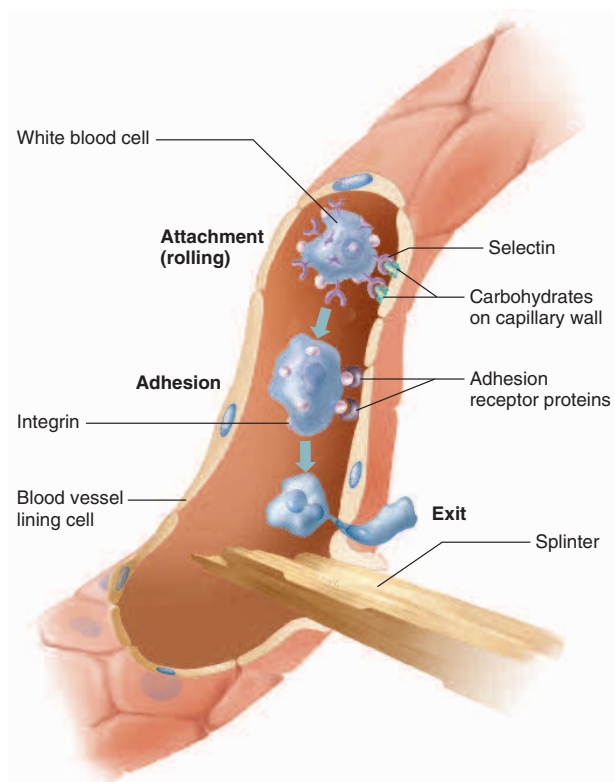


Figure 2.21 Cellular adhesion. Cellular adhesion molecules (CAMs), including selectins, integrins, and adhesion receptor proteins, direct white blood cells to injury sites.

If the signals that direct white blood cells to injury sites fail, a condition called leukocyte-adhesion deficiency (MIM 116920) results. The first symptom is often teething sores that do not heal. These and other small wounds never accumulate the pus (bacteria, cellular debris, and white blood cells) that indicates the body is fighting infection. The person lacks the CAMs that enable white blood cells to stick to blood vessel walls, and so blood cells travel right past wounds. An affected individual must avoid injury and infection, and receive anti-infective treatments for even the slightest wound.

More common disorders may also reflect abnormal cellular adhesion. Cancer cells journey easily from one part of the body to another thanks to impaired cellular adhesion. Arthritis may occur when the wrong adhesion molecules rein in white blood cells, inflaming a joint where no injury exists.

Cellular adhesion is critical to many other functions. CAMs guide cells surrounding an embryo to grow toward maternal cells and form the placenta, the supportive organ linking a pregnant woman to the fetus. Sequences of CAMs also help establish connections among nerve cells in the brain that underlie learning and memory.

Key Concepts

1. In signal transduction, cell surface receptors receive information from first messengers (stimuli) and pass them to second messengers, which then trigger a cellular response.
2. Cellular adhesion molecules (CAMs) guide white blood cells to injury sites using a sequence of cell-protein interactions.

2.5 Stem Cells

Bodies grow and heal thanks to cells that retain the ability to divide, generating both new cells like themselves and cells that go on to specialize. **Stem cells** and **progenitor cells** renew tissues so that as the body grows, or loses cells to apoptosis, injury, and disease, other cells are produced that take their places.

Cell Lineages

A stem cell divides by mitosis to yield either two daughter cells that are stem cells like itself, or one that is a stem cell and one that is a partially specialized progenitor cell (**figure 2.22**). The characteristic of **self-renewal** is what makes a stem cell a stem cell—its ability to continue the lineage of cells that can divide to give rise to another cell like itself. A progenitor cell cannot self-renew, and its daughters specialize as any of a restricted number of cell types. A fully differentiated cell, such as a mature blood cell, descends from a sequence of increasingly specialized progenitor cell intermediates, each one less like a stem cell and more like a blood cell. Our more than 260 differ-

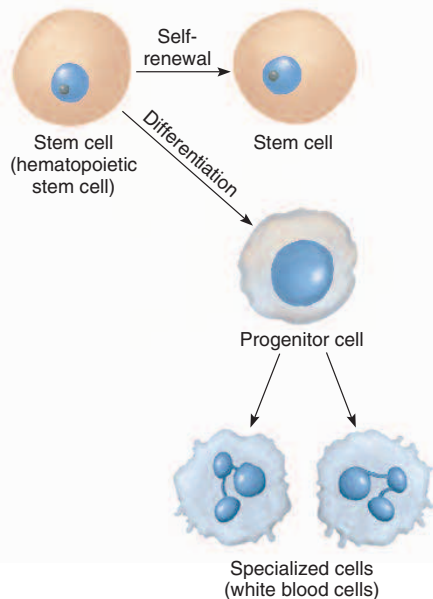


Figure 2.22 Stem cells and progenitor cells. A stem cell is less specialized than the progenitor cell that descends from it by mitosis. Various types of stem cells provide the raw material for producing the specialized cells that comprise tissues, while retaining the ability to generate new cells. A hematopoietic stem cell resides in the bone marrow and can produce progenitors whose daughter cells may specialize as certain blood cell types.

entiated cell types develop from lineages of stem and progenitor cells. **Figure 2.23** shows parts of a few lineages.

Stem cells and progenitor cells are described in terms of developmental potential—that is, according to the number of possible fates of their daughter cells. A fertilized ovum is the ultimate stem cell. It is totipotent, which means that it can give rise to every cell type, including the cells of the membranes that support the embryo. Other stem cells and progenitor cells are pluripotent: Their daughter cells have fewer possible fates. Some are multipotent: Their daughter cells have only a few developmental “choices.” This is a little like a freshman’s consideration of many majors, compared to a junior’s more narrowed focus in selecting courses.

As stem cell descendants specialize, they express some genes and ignore others. An immature bone cell forms from a progenitor cell by manufacturing mineral-binding proteins and enzymes. In contrast, an immature muscle cell forms from a muscle progenitor cell that accumulates contractile proteins. The bone cell does not produce muscle proteins, nor does the muscle cell produce bone proteins. All cells, however, synthesize proteins for basic “housekeeping” functions, such as energy acquisition and protein synthesis.

Many, if not all, of the organs in an adult human body have stem or progenitor cells. These cells can divide when injury or illness occurs and generate new cells to replace

damaged ones. Stem cells in the adult may have been set aside in the embryo or fetus in particular organs as repositories of future healing. Evidence suggests that some stem cells, such as those from bone marrow, can travel to and replace damaged or dead cells elsewhere in the body, in response to signals that are released in injury or disease. Because every cell contains all of an individual’s genetic material, any cell type, given appropriate signals, can in theory become any other. This concept is the basis of much of stem cell technology.

Stem Cells in Health Care

Researchers are tapping into the natural versatility of stem cells to learn more about basic biology and to develop treatments for a great variety of diseases and injuries—not just inherited conditions. Thousands of clinical trials are currently testing stem cell-based treatments. These cells come from donors as well as from patients’ own bodies, as **figure 2.24** illustrates. The cells can be mass-produced in laboratory glassware, and if they originate with a patient’s cell or nucleus, they are a genetic match.

Stem Cell Sources

There are three general sources of human stem cells.

Embryonic stem (ES) cells are not actually cells from an embryo, but are created in a laboratory dish using certain cells from a region of a very early embryo called an inner cell mass (ICM). Some ICM cells, under certain conditions, become pluripotent and can self-renew—they are stem cells. The ICM cells used to derive ES cells can come from two sources: “left-over” embryos from fertility clinics that would otherwise be destroyed, and from nuclear transfer, in which a nucleus from a person’s somatic cell is transferred to an egg cell that has had its own nucleus removed. Nuclear transfer is popularly called “cloning” because it copies the person’s genome.

Induced pluripotent stem (iPS) cells are somatic cells that are “reprogrammed” to differentiate into any of several cell types. This change may require a journey back through developmental time to an ES cell-like state, then to specialize anew as a different, desired cell type. Or, cells can be reprogrammed directly into another cell type. The instructions for a cell to reprogram are sent into cells as genes—four in particular that seem to have the ability to alter a cell’s fate. These factors can be delivered in various ways. Deriving iPS cells does not require the use of any cells from an embryo. However, researchers do not yet know whether iPS cells function exactly as ES cells. For research purposes, human ES cells remain the “gold standard” because they more closely approximate normal development than do iPS cells. Once we learn more about how accurately iPS cells mimic normal development, human ES cells might no longer be required.

To return to the college major analogy, the idea of reprogramming a cell is like a senior in college deciding to change his or her major. If, for example, a French major wanted to become a mechanical engineer, he’d have to start over, taking very different courses. But if a biology major wanted to become

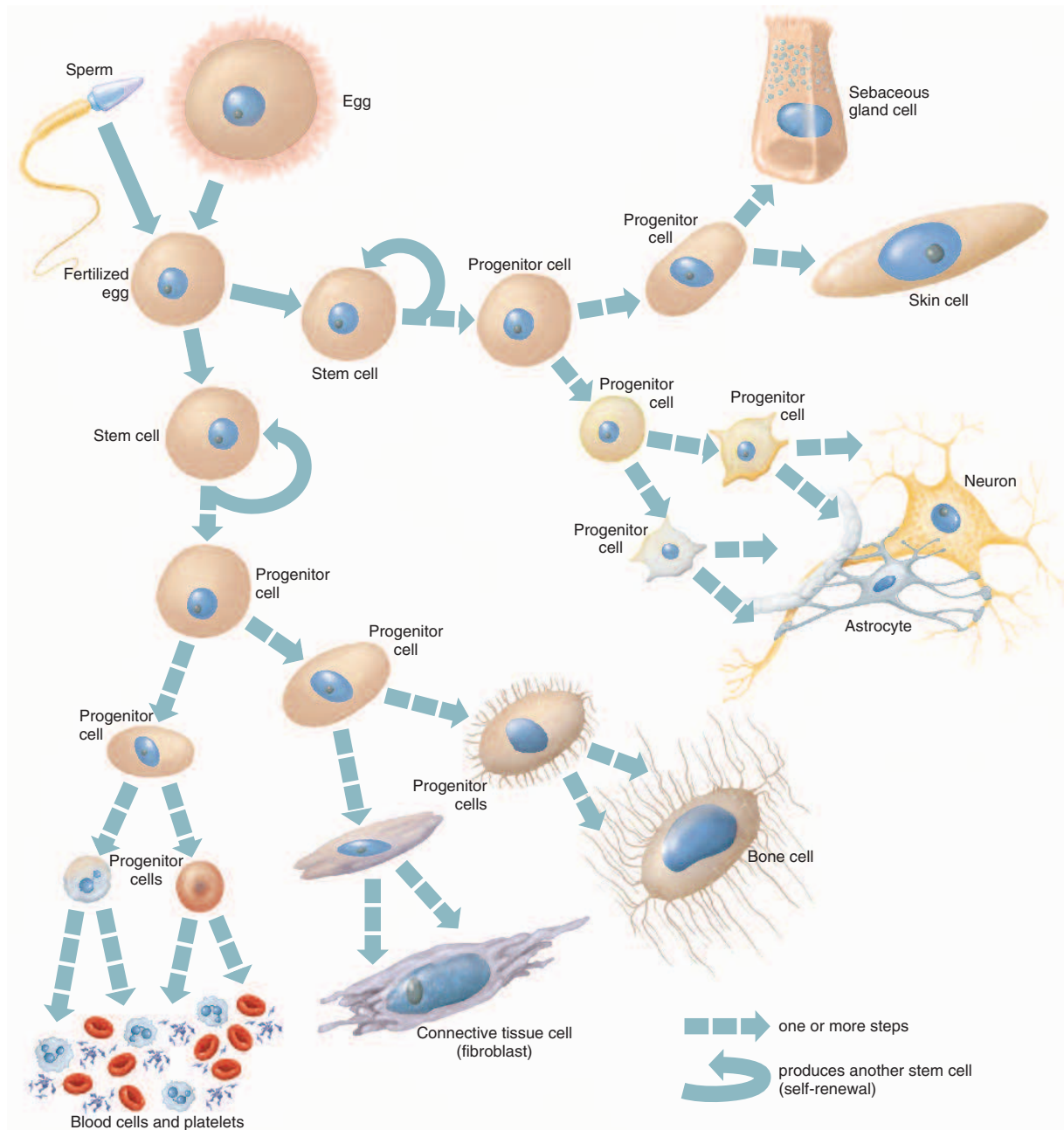


Figure 2.23 Pathways to cell specialization. All cells in the human body descend from stem cells, through the processes of mitosis and differentiation. The differentiated cells on the lower left are all connective tissues (blood, connective tissue, and bone), but the blood cells are more closely related to each other than they are to the other two cell types. On the upper right, the skin and sebaceous gland cells share a recent progenitor, and both share a more distant progenitor with neurons and supportive astrocytes. Imagine how complex the illustration would be if it embraced all 260-plus types of cells in a human body!

a chemistry major, she would not need to start from scratch because many of the same courses apply to both majors. So it is for stem cells. Taking a skin cell from a man with heart disease and turning it into a healthy heart muscle cell might require

taking that initial cell back to an ES or iPS state, because these cells come from very different lineages. But turning an endocrine cell of the pancreas into a digestive cell of the pancreas requires fewer steps backwards.

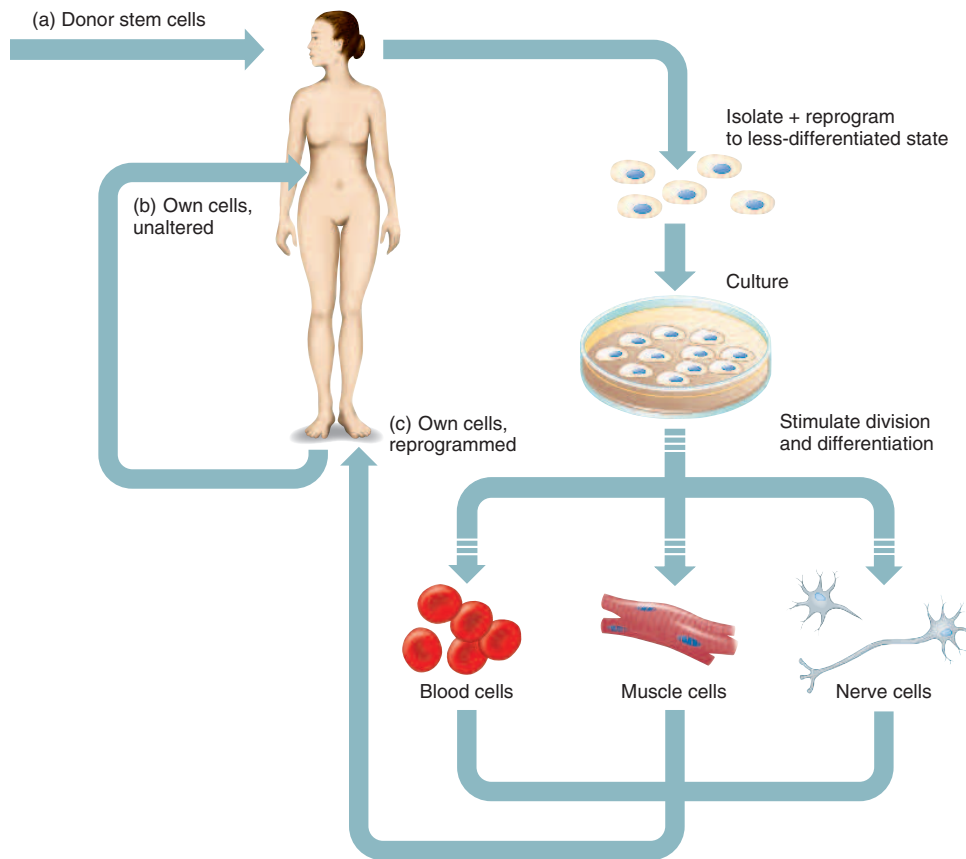


Figure 2.24 Using stem cells to heal. (a) Stem cells from donors (bone marrow or umbilical cord blood) are already in use. (b) A person’s cells may be used, unaltered, to replace damaged tissue, such as bone marrow. (c) It is possible to “reprogram” a person’s cells in culture, taking them back to a less specialized state and then nurturing them to differentiate as a needed cell type.

“Adult” stem cells are more accurately called tissue-specific or somatic stem cells because they are found in the tissues of fetuses, embryos and children, and not just in adult bodies. Adult stem cells self-renew, but most are multipotent, giving rise to a few types of specialized daughter cells. Researchers are still discovering niches of adult stem cells in the body. Many potentially valuable adult stem cells are routinely discarded as medical waste.

Stem Cell Applications

Stem cells are being used in four basic ways. In drug discovery and development, stem cell cultures supply the human cells that are affected in a particular disease, which may be difficult or impossible to culture. Drugs are tested on these cells. Liver and heart cells derived from stem cells are particularly useful in testing drugs for side effects, because the liver detoxifies many drugs, and the heart is easily harmed by them. Using stem cells

in drug development can minimize the need to experiment on animals and can weed out drugs with adverse effects before they are tested on people.

A second application of stem cells growing in culture is to observe the earliest signs of a disease, which may begin long before symptoms appear in a person. The first disease for which human iPS cells were derived was amyotrophic lateral sclerosis (ALS), also known as Lou Gehrig’s disease (**figure 2.25**). In ALS, motor neurons that enable a person to move gradually fail. A few years after the first signs of weakness or stumbling, death comes from failure of the respiratory muscles. ALS has been difficult to study because motor neurons cannot be cultured in the laboratory because they do not divide. However, iPS cells derived from fibroblasts in patients’ skin can be reprogrammed in culture to become ALS motor neurons. Thanks to these and other iPS cells, researchers are now observing the beginnings of hundreds of diseases—and discovering new ways to treat them.

The third application of stem cells is in implants and transplants, for treatments. This approach is not new—the

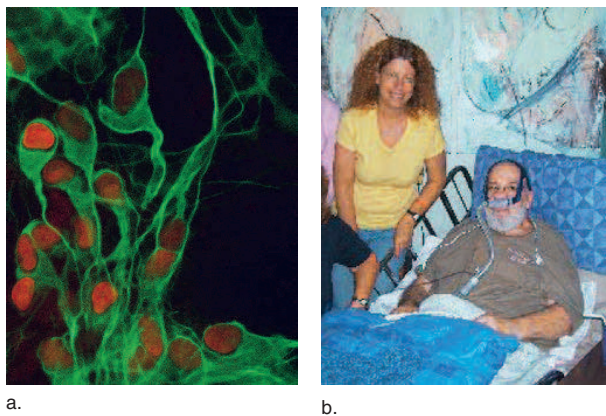


Figure 2.25 (a) Reprogrammed motor neurons from an ALS patient enable researchers to study the beginnings of the disease. (b) Glenn Nichols was an English teacher and high school principal who lived with and died of ALS. The author, in yellow, was his hospice volunteer. This book is dedicated to him.

Bioethics: Choices for the Future

Should You Bank Your Stem Cells?

The parents-to-be were very excited by the DVD that came in the mail shortly after they began seeing an obstetrician:

“Bank your baby’s cord blood stem cells and benefit from breakthroughs. Be prepared for the unknowns in life.”

The short film profiled children who were saved from deadly diseases because their parents had stored their umbilical cord blood. The statistics quoted were persuasive: More than 70 diseases are currently treatable with cord blood transplants, and 10,000 procedures have already been done.

With testimonials like that, it is little wonder that parents collectively spend more than \$100 million per year to store cord blood. The ads and statistics are indeed accurate, but they are misleading, because of what they *don’t* say: Most people never actually use the umbilical cord blood stem cells that they store. The scientific reasons go beyond the fact that the diseases that are treatable so far are very rare. In addition, cord blood stem cells are not nearly as pluripotent as some other stem cells, limiting their applicability. Perhaps the most compelling reason that stem cell banks are rarely used is based on logic: For a person with a blood disorder or an inherited disease, *healthy* stem cells are required—not his or her own, which could cause the disease all over again.

Add to these caveats the cost of a commercial cord blood bank—typically more than \$1,000 for the initial collection plus an annual fee—and it is clear that a not-for-profit bank would help more people. Cord blood stem cells are donated to these banks not to help the individuals who donate them, but whoever can use them. The U.S. National Institutes of Health and many other nations have supported such not-for-profit banks for years.

As stem cell science has leaped forward, however, both commercial cell banks and anecdotal reports of successes have captured much media attention. This was the case for 18-month-old Dallas Hextell. His cerebral palsy greatly improved after he was treated

with his own cord blood cells, but whether he would have improved without the treatment isn’t known.

Banking on the likelihood that ongoing clinical trials, such as the one that helped young Dallas, will ultimately demonstrate great value in autologous (“self”) stem cell treatments, other companies have entered the marketplace—and not just for newborns. One company, for example, offers to bank “very small embryonic-like stem cells” for an initial charge of \$7,500 and a \$750 annual fee, “enabling people to donate and store their own stem cells when they are young and healthy for their personal use in times of future medical need.” The cells come from a person’s blood and, in fact, one day may be very useful, but the research has yet to be done supporting use of the cells in treatments.

Questions for Discussion

1. Storing stem cells is not regulated by the U.S. government the way that a drug or a surgical procedure is because it is a service that will be helpful for treatments not yet invented. Do you think such banks should be regulated, and if so, by whom and how?
2. What information do you think that companies offering to store stem cells should present on their websites?
3. Do you think that advertisements for cord blood storage services that have quotes and anecdotal reports, but do not mention that most people who receive stem cell transplants do not in fact receive their own cells, are deceptive? Or do you think it is the responsibility of the consumer to research and discover this information?
4. How can medical consumers become aware that the government funds facilities to store stem cells?
5. It is highly likely that in the future, many stem cell-based treatments will indeed be possible, following large-scale clinical trials. What is the fairest way to prepare for this type of future medical treatment?

oldest such treatment, a bone marrow transplant, has been around for more than half a century. Many other uses of adult stem cells, delivered as implants, transplants, or simply infusions into the bloodstream, are being tested. A patient’s own bone marrow cells, for example, can be removed, bathed in selected factors, and reinfused. They follow signals that lead them to damaged tissues.

The fourth application of stem cells became clear with the creation of iPS cells. It might be possible to introduce the reprogramming proteins directly into the body to stimulate stem cells in their natural niches. Once we understand the signals, we might not need the cells. The applications of stem cells seem limited only by our imaginations. *Bioethics: Choices for the Future* discusses stem cell banking.

Key Concepts

1. A stem cell self-renews and gives rise to any of several differentiated cell types. All cells descend from stem cells and progenitor cells, which do not self-renew.
2. Cells differentiate down cell lineages by differential gene expression.
3. Stem cells are present throughout life and provide growth and repair.
4. Stem cells in health care include embryonic stem cells, induced pluripotent stem cells, and adult stem cells. ES and reprogrammed cells give researchers views of disease origins.

Summary

2.1 Introducing cells

1. Cells are the fundamental units of life and comprise the human body. Inherited traits and illnesses can be understood at the cellular and molecular levels.
2. All cells share certain features, but they are also specialized because they express different subsets of genes.
3. **Somatic** (body) cells are **diploid** and sperm and egg cells are **haploid**. **Stem cells** produce new cells.

2.2 Cell Components

4. Prokaryotic cells are small and lack **nuclei** and other **organelles**. **Eukaryotic** cells have organelles, and their DNA is in a **nucleus**.
5. Cells consist primarily of water and several types of macromolecules: **carbohydrates, lipids, proteins, and nucleic acids**.
6. Organelles sequester related biochemical reactions, improving the efficiency of life functions and protecting the cell. The cell also consists of **cytoplasm** and other chemicals.
7. The nucleus contains DNA and a nucleolus, which is a site of ribosome synthesis. **Ribosomes** provide scaffolds for protein synthesis; they exist free in the cytoplasm or complexed with the **rough endoplasmic reticulum** (ER).
8. In secretion, the rough ER is the site of protein synthesis and folding, the **smooth ER** is the site of lipid synthesis, transport, and packaging, and the **Golgi apparatus** packages secretions into vesicles, which exit through the **plasma membrane**. **Lysosomes** contain enzymes that dismantle debris, and **peroxisomes** house enzymes that perform a variety of functions. Enzymes in **mitochondria** extract energy from nutrients.
9. The plasma membrane is a protein-studded phospholipid bilayer. It controls which substances exit and enter the cell, and how the cell interacts with other cells.
10. The **cytoskeleton** is a protein framework of hollow microtubules, made of tubulin, and solid microfilaments, which consist of actin. Intermediate filaments are made of more than one protein type and are abundant in skin. The cytoskeleton and the plasma membrane distinguish different types of cells.

2.3 Cell Division and Death

11. Coordination of cell division (**mitosis**) and cell death (**apoptosis**) maintains cell numbers, enabling structures to enlarge during growth and development but preventing abnormal growth.

12. The **cell cycle** describes whether a cell is dividing (mitosis) or not (**interphase**). Interphase consists of two gap phases, when proteins and lipids are produced, and a synthesis phase, when DNA is replicated.
13. Mitosis proceeds in four stages. In **prophase**, replicated chromosomes consisting of two **chromatids** condense, the **spindle** assembles, the nuclear membrane breaks down, and the nucleolus is no longer visible. In **metaphase**, replicated chromosomes align along the center of the cell. In **anaphase**, the **centromeres** part, equally dividing the now unreplicated chromosomes into two daughter cells. In **telophase**, the new cells separate. Cytokinesis apportions other components into daughter cells.
14. Internal and external factors control the cell cycle. Checkpoints are times when proteins regulate the cell cycle. **Telomere** (chromosome tip) length determines how many more mitoses will occur. Crowding, hormones, and growth factors signal cells from the outside; the interactions of cyclins and kinases trigger mitosis from inside.
15. In apoptosis, a receptor on the plasma membrane receives a death signal, which activates caspases that tear apart the cell in an orderly fashion. Membrane surrounds the pieces, preventing inflammation.

2.4 Cell-Cell Interactions

16. In **signal transduction**, a stimulus (first messenger) activates a cascade of action among membrane proteins, culminating in the production of a second messenger that turns on enzymes that provide the response.
17. **Cellular adhesion** molecules enable cells to interact. Selectins slow the movement of leukocytes, and integrins and adhesion receptor proteins guide the blood cell through a capillary wall to an injury site.

2.5 Stem Cells

18. **Stem cells** produce daughter cells that retain the ability to divide and daughter cells that specialize. Progenitor cells give rise to more specialized daughter cells but do not self-renew.
19. A fertilized ovum is totipotent. Some stem cells are pluripotent, and some are multipotent. Cells are connected in lineages.
20. The three sources of stem cells are **embryonic stem cells, induced pluripotent stem cells**, and adult stem cells.
21. Stem cell technology enables researchers to observe the origins of diseases and to devise new types of treatments.

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Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

- Match each organelle to its function.

Organelle	Function
a. lysosome	1. lipid synthesis
b. rough ER	2. houses DNA
c. nucleus	3. energy extraction
d. smooth ER	4. dismantles debris
e. Golgi apparatus	5. detoxification
f. mitochondrion	6. protein synthesis
g. peroxisome	7. processes secretions
- What advantage does compartmentalization provide to a large and complex cell?
- Explain the functions of the following proteins:
 - tubulin and actin
 - caspases
 - cyclins and kinases
 - checkpoint proteins
 - cellular adhesion molecules
- List four types of controls on cell cycle rate.
- How can all of a person's cells contain exactly the same genetic material, yet specialize as bone cells, nerve cells, muscle cells, and connective tissue cells?
- Distinguish between
 - a bacterial cell and a eukaryotic cell.
 - interphase and mitosis.
 - mitosis and apoptosis.
 - rough ER and smooth ER.
 - microtubules and microfilaments.
 - a stem cell and a progenitor cell.
 - totipotent and pluripotent.
- Select a process described in the chapter (such as signal transduction or apoptosis). List the steps and state why the cell could not survive without this ability.
- How are intermediate filaments similar to microtubules and microfilaments, and how are they different?
- What role does the plasma membrane play in signal transduction?
- Distinguish among ES cells, iPS cells, and adult stem cells, and state the pros and cons of working with each to develop a therapy.

Applied Questions

- How might abnormalities in each of the following contribute to cancer?
 - cellular adhesion
 - signal transduction
 - balance between mitosis and apoptosis
 - cell cycle control
 - telomerase activity
- How do stem cells maintain their populations within tissues that consist of mostly differentiated cells?
- Why wouldn't a cell in an embryo likely be in phase G_0 ?
- A defect in which organelle would cause fatigue?
- If you wanted to create a synthetic organelle to test new drugs for toxicity, which natural organelle's function would you try to replicate?
- An inherited form of migraine is caused by a mutation in a gene (*SCN1A*) that encodes a sodium channel in neurons. What is a sodium channel, and in which cell structure is it located?
- How can signal transduction, the plasma membrane, and the cytoskeleton function together?
- A single stem cell in skin gives rise to skin cells, hair follicle cells, and sebaceous (oil) gland cells. Suggest a treatment that might use these cells.
- Invent a stem cell therapy. Choose a disease, identify the affected cell types, describe how they are abnormal, and explain the type of stem cell (ES, iPS, or adult) you would use. Explain how the treatment would work, and what the dangers might be.
- Ads proclaiming "*Stem Cells: The Future of Skin Rejuvenation*," have appeared in many magazines. The product is actually secretions from stressed, cultured skin cells from behind the ears of healthy young people. The secretions are dried out and mixed with conventional skin cream. The ad promises that the goop contains "signals delivered by skin stem cells," because skin has stem cells.
 - Is this product worth \$155 for a 1.7-ounce jar? Why or why not?
 - What evidence might convince you to buy such a product?

Web Activities

- Several companies offer expensive "banking" of stem cells, even if treatments using the cells have not yet been invented. Find a website offering to bank the cells, and discuss whether or not the company provides enough information for you to make an informed choice as to whether or not to use the service.
- Nations vary greatly in which stem cell technologies they consider ethical. Go to www.hinxongroup.org/wp.html and use the interactive map feature to find examples of
 - a nation that allows use of human embryos from clinics and nuclear transfer
 - a nation that allows use of clinic embryos but not nuclear transfer
 - a nation that bans derivation of any new human ES cell lines

Case Studies and Research Results

13. Nadine has a form of neuronal ceroid lipofuscinosis (MIM 610127). She suffers from seizures, loss of vision, and lack of coordination, and will likely not survive beyond 10 years of age. Her cells lack an enzyme that normally breaks down certain proteins, causing them to accumulate and destroy the nervous system. Name two organelles that are affected in this illness.
14. Studies show that women experiencing chronic stress, such as from caring for a severely disabled child, have telomeres that shorten at an accelerated rate. Suggest a study that would address the question of whether men have a similar reaction to chronic stress.

Forensics Focus

15. Michael Mastromarino was recently sentenced to serve many years in prison for trafficking in body parts. As the owner of Biomedical Tissue Services in Fort Lee, NJ, Mastromarino and his cohorts dismembered corpses taken, without any consent, from funeral homes in Pennsylvania, New Jersey, and New York. (An unwitting donor was Alistair Cooke, the host of “Masterpiece Theatre” on television.) Thousands of parts from hundreds of bodies were used in various surgical procedures, including hip and knee replacements and dental implants. The most commonly used product was a paste made of bone. Many family members testified at the trials. What is the basis of the fact that cells from bone tissue can be matched to blood or cheek lining cells from blood relatives?

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CHAPTER

3

Meiosis and Development

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Selling Eggs: Vanessa's Story

"I couldn't believe the ad in the student newspaper—a semester's tuition for a few weeks of discomfort! So I applied. I was 18, on the volleyball team, healthy except for some acne, and had a 3.8 GPA. Since I didn't plan on having children at the time, or at all, I thought why not?"

I passed the physical and psychological screens, and my family history seemed OK. I was accepted! Then three weeks later, I got the call. A young couple who couldn't have a child because the woman had had cancer wanted to use donor eggs, to be fertilized *in vitro* by the man's sperm. They'd seen my photo and read my file, and thought I'd be a good match. I was thrilled, but the warnings scared me: bleeding, infection, cramping, mood swings, and scarred ovaries.

For the first 10 days, I gave myself shots in the thigh of a drug to suppress my ovaries. Then for the next 12 days, I injected myself with two other drugs in the back of the hip, to mature my egg cells. Frequent ultrasounds showed that my ovaries looked like grape clusters, with the maturing eggs popping to the surface. Towards the end of the regimen I felt a dull aching in my belly.

The egg retrieval wasn't bad. I was sedated, had anesthesia, and the doctor removed 20 eggs using a needle passed through the wall of my vagina. My abdomen ached at night and the next day, and I felt bloated for a few days. But they got a dozen eggs! Two were implanted, four frozen, and the rest donated for stem cell research.

That was 4 years ago. One of my eggs may now be a preschooler!"

3.1 The Reproductive System

Genes orchestrate our physiology from a few days after conception through adulthood. Expression of specific sets of genes sculpts the differentiated cells that interact, aggregate, and fold, forming the organs of the body. Abnormal gene functioning can affect health at all stages of development. Certain single-gene mutations act before birth, causing broken bones, dwarfism, or even cancer. Many other mutant genes exert their effects during childhood. It may take parents months or even years to realize their child has a health problem. Inherited forms of heart disease and breast cancer can appear in early or middle adulthood, which is earlier than multifactorial forms of these conditions. Pattern baldness is a very common single-gene trait that may not become obvious until well into adulthood.

This chapter explores the stages of the human life cycle that form the backdrop against which genes function.

The first cell that leads to development of a new individual forms when a **sperm** from a male and an **oocyte** (also called an egg) from a female join. Sperm and oocytes are **gametes**, or sex cells. They provide a mechanism for mixing genetic material from past generations. Because we have thousands of genes, some with many variants, each person (except for identical twins) has a unique combination of inherited traits.

Sperm and oocytes are produced in the reproductive system. The reproductive organs are organized similarly in the male and female. Each system has

- paired structures, called **gonads**, where the sperm and oocytes are manufactured;
- tubular structures that transport these cells; and
- hormones and secretions that control reproduction.

The Male

Sperm cells develop within a 125-meter-long network of seminiferous tubules, which are packed into paired, oval organs called **testes** (sometimes called testicles) (**figure 3.1**). The testes are the male gonads. They lie outside the abdomen within a sac called the scrotum. This location keeps the testes cooler than the rest of the body, which is necessary for sperm to develop. Leading from each testis is a tightly coiled tube, the epididymis, in which sperm cells mature and are stored. Each epididymis continues into another tube, the ductus deferens. Each ductus deferens bends behind the bladder and joins the urethra, which is the tube that carries sperm and urine out through the penis.

Along the sperm's path, three glands add secretions. The ductus deferentia pass through the prostate

gland, which produces a thin, milky, alkaline fluid that activates the sperm to swim. Opening into the ductus deferens is a duct from the seminal vesicles, which secrete fructose (an energy-rich sugar) and hormonelike prostaglandins, which may stimulate contractions in the female that help sperm and oocyte meet. The bulbourethral glands, each about the size of a pea, join the urethra where it passes through the body wall. They secrete an alkaline mucus that coats the urethra before sperm are released. All of these secretions combine to form the seminal fluid that carries sperm.

During sexual arousal, the penis becomes erect so that it can penetrate and deposit sperm in the female reproductive tract. At the peak of sexual stimulation, a pleasurable sensation called orgasm occurs, accompanied by rhythmic muscular contractions that eject the sperm from each ductus deferens through the urethra and out the penis. The discharge of sperm from the penis, called ejaculation, delivers about 200 to 600 million sperm cells.

The Female

The female sex cells develop in paired organs in the abdomen called **ovaries** (**figure 3.2**), which are the female gonads. Within each ovary of a newborn girl are about a million immature oocytes. Each individual oocyte nestles within nourishing follicle cells, and each ovary houses oocytes in different stages of development. After puberty, about once a month, one ovary releases the most mature oocyte. Beating cilia sweep the mature oocyte into the fingerlike projections of one of two uterine (also called fallopian) tubes. The tube carries the oocyte into a muscular, saclike organ called the uterus, or womb.

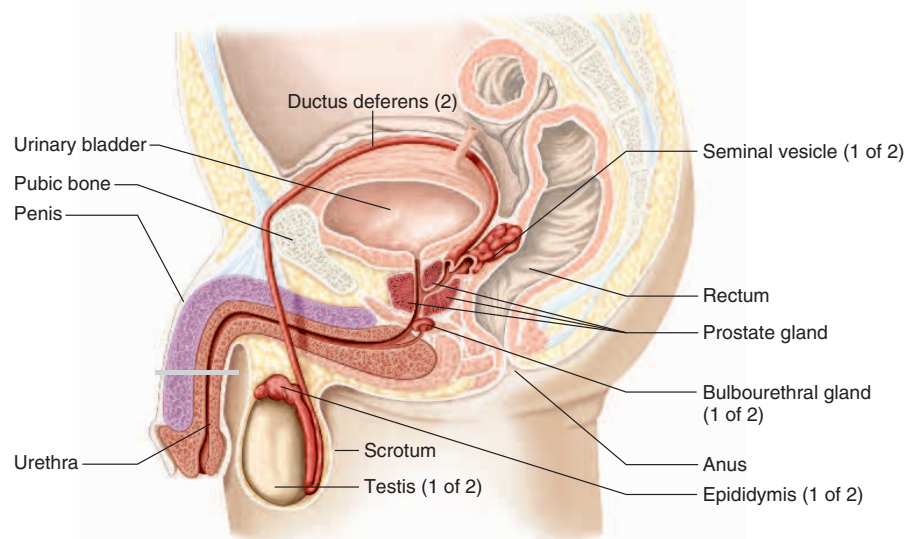


Figure 3.1 The human male reproductive system. Sperm cells are manufactured in the seminiferous tubules, which wind tightly within the testes, which descend into the scrotum. The prostate gland, seminal vesicles, and bulbourethral glands add secretions to the sperm cells to form seminal fluid. Sperm mature and are stored in the epididymis and exit through the ductus deferens. The paired ductus deferentia join in the urethra, which transports seminal fluid from the body.

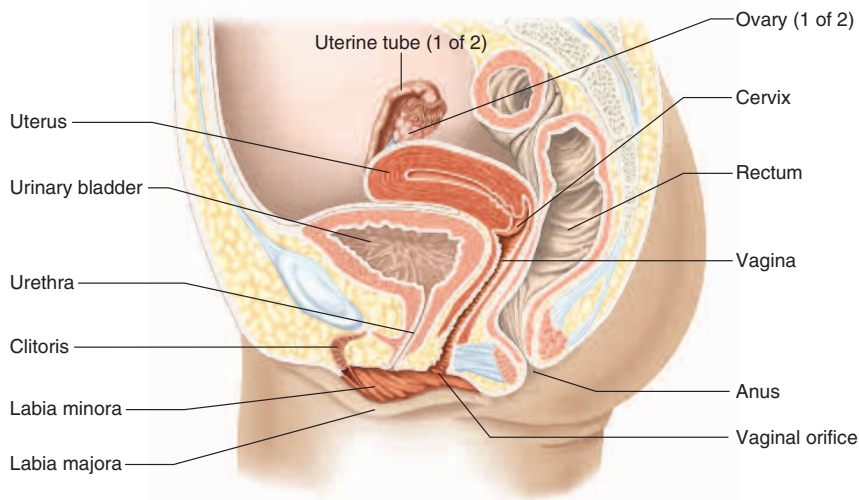


Figure 3.2 The human female reproductive system. Oocytes mature in the paired ovaries. Once a month after puberty, an ovary releases one oocyte, which is drawn into a nearby uterine tube. If a sperm fertilizes the oocyte in the uterine tube, the fertilized ovum continues into the uterus, where for nine months it divides and develops. If the oocyte is not fertilized, the body expels it, along with the built-up uterine lining. This is the menstrual flow.

The released oocyte may encounter a sperm. This usually occurs in a uterine tube. If the sperm enters the oocyte and the DNA of the two gametes merges into a new nucleus, the result is a fertilized ovum. After about a day, this first cell divides while moving through the uterine tube. It then settles into the lining of the uterus, where it may continue to divide and an embryo develops. If fertilization does not occur, the oocyte, along with much of the uterine lining, is shed as the menstrual flow. Hormones coordinate the monthly menstrual cycle.

The lower end of the uterus narrows and leads to the cervix, which opens into the tubelike vagina. The vaginal opening is protected on the outside by two pairs of fleshy folds. At the upper juncture of both pairs is the 2-centimeter-long clitoris, which is anatomically similar to the penis. Rubbing the clitoris triggers female orgasm. Hormones control the cycle of oocyte maturation and the preparation of the uterus to nurture a fertilized ovum.

Key Concepts

1. Sperm develop in the seminiferous tubules, mature and collect in each epididymis, enter the ductus deferentia, and move through the urethra in the penis. The prostate gland adds an alkaline fluid, seminal vesicles add fructose and prostaglandins, and bulbourethral glands secrete mucus to form seminal fluid.
2. In the female, ovaries contain oocytes. Each month, an ovary releases an oocyte, which enters a uterine tube leading to the uterus. If the oocyte is fertilized, it begins rapid cell division and nestles into the uterine lining to divide and develop. Otherwise, the oocyte exits the body. Hormones control the cycle of oocyte development.

3.2 Meiosis

Gametes form from special cells, called germline cells, in a type of cell division called **meiosis** that halves the chromosome number. A further process, maturation, sculpts the distinctive characteristics of sperm and oocyte. The organelle-packed oocyte has 90,000 times the volume of the streamlined sperm.

Gametes contain 23 different chromosomes—half the usual amount of genetic material, but still a complete genome. In contrast, somatic cells contain 23 pairs, or 46 chromosomes. One member of each pair comes from the person's mother and one comes from the father. The chromosome pairs are called **homologous pairs**, or *homologs* for short. Homologs have the same genes in the same order but may carry different alleles, or variants, of the same gene. Recall from chapter 2 that gametes are haploid ($1n$), which means that they have

only one of each type of chromosome, and somatic cells are diploid ($2n$), with two copies of each chromosome type.

Halving the number of chromosomes in gametes makes sense. If the sperm and oocyte each contained 46 chromosomes, the fertilized ovum would have twice the normal number of chromosomes, or 92. Such a genetically overloaded cell, called a polyploid, usually does not develop normally, or at all. About one in a million newborns has polyploid cells, and has abnormalities in all organ systems and usually only lives a few days. Studies on spontaneously aborted embryos indicate that about 1 percent of conceptions have three chromosome sets instead of the normal two. Therefore, most polyploid embryos do not survive to be born. Other chromosome abnormalities, such as a missing or extra chromosome, are more common than polyploidy.

In addition to producing gametes, meiosis mixes up trait combinations. For example, a person might produce one gamete containing alleles encoding green eyes and freckles, yet another gamete with alleles encoding brown eyes and no freckles. Meiosis explains why siblings differ genetically from each other and from their parents.

In a much broader sense, meiosis, as the mechanism of sexual reproduction, provides genetic diversity, which enables a population to survive a challenging environmental change. A population of sexually reproducing organisms is made up of individuals with different genotypes and phenotypes. In contrast, a population of asexually reproducing organisms such as bacteria, or genetically identical crops, consists of individuals with the same genome sequence. Should a new threat arise, such as an infectious disease that kills only organisms with a certain genotype, then the entire asexual population could be wiped out. However, in a sexually reproducing population, individuals that inherited a certain combination of genes might survive. This

differential survival of certain genotypes is the basis of evolution, discussed in chapter 16.

Meiosis entails two divisions of the genetic material. The first division is called a **reduction division** (or meiosis I) because it reduces the number of replicated chromosomes from 46 to 23. The second division, called an **equational division** (or meiosis II), produces four cells from the two cells formed in the first division by splitting the replicated chromosomes. **Figure 3.3** shows an overview of the process, and **figure 3.4** depicts the major events of each stage.

As in mitosis, meiosis occurs after an interphase period when DNA is replicated (doubled) (**table 3.1**). For each chromosome pair in the cell undergoing meiosis, one homolog comes from the person's mother, and one from the father. In figures 3.3 and 3.4, the colors represent the contributions of the two parents, whereas size indicates different chromosomes.

After interphase, prophase I (so called because it is the prophase of meiosis I) begins as the replicated chromosomes condense and become visible when stained. A spindle forms. Toward the middle of prophase I, the homologs line up next to one another, gene by gene, in an event called **synapsis**. A mixture of RNA and protein holds the chromosome pairs together. At this time, the homologs exchange parts in a process called **crossing over** (**figure 3.5**). All four chromatids that comprise each homologous chromosome pair are pressed together as exchanges

occur. After crossing over, each homolog bears some genes from each parent. Prior to this, all of the genes on a homolog were derived from one parent. New gene combinations arise from crossing over when the parents carry different alleles. Toward the end of prophase I, the synapsed chromosomes separate but remain attached at a few points along their lengths.

To understand how crossing over mixes trait combinations, consider a simplified example. Suppose that homologs carry genes for hair color, eye color, and finger length. One of the chromosomes carries alleles for blond hair, blue eyes, and short fingers. Its homolog carries alleles for black hair, brown eyes, and long

Table 3.1 Comparison of Mitosis and Meiosis

Mitosis	Meiosis
One division	Two divisions
Two daughter cells per cycle	Four daughter cells per cycle
Daughter cells genetically identical	Daughter cells genetically different
Chromosome number of daughter cells same as that of parent cell ($2n$)	Chromosome number of daughter cells half that of parent cell ($1n$)
Occurs in somatic cells	Occurs in germline cells
Occurs throughout life cycle	In humans, completes after sexual maturity
Used for growth, repair, and asexual reproduction	Used for sexual reproduction, producing new gene combinations

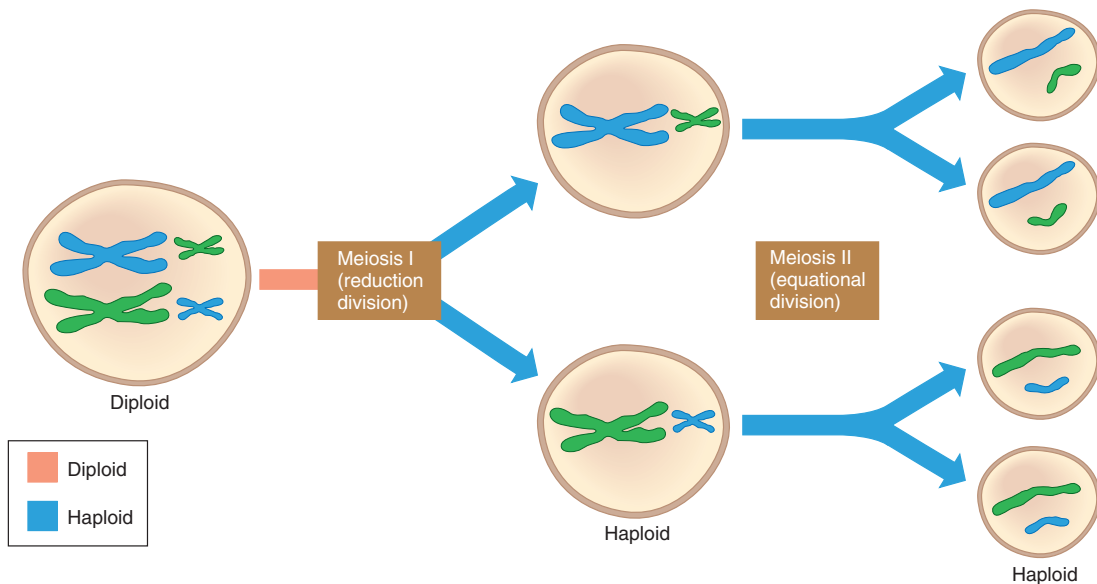


Figure 3.3 Overview of meiosis. Meiosis is a form of cell division in which certain cells are set aside and give rise to haploid gametes. This simplified illustration follows the fate of two chromosome pairs rather than the true 23 pairs. In actuality, the first meiotic division reduces the number of chromosomes to 23, all in the replicated form. In the second meiotic division, the cells essentially undergo mitosis. The result of the two meiotic divisions (in this illustration and in reality) is four haploid cells. In this illustration, homologous pairs of chromosomes are indicated by size, and parental origin of chromosomes by color.

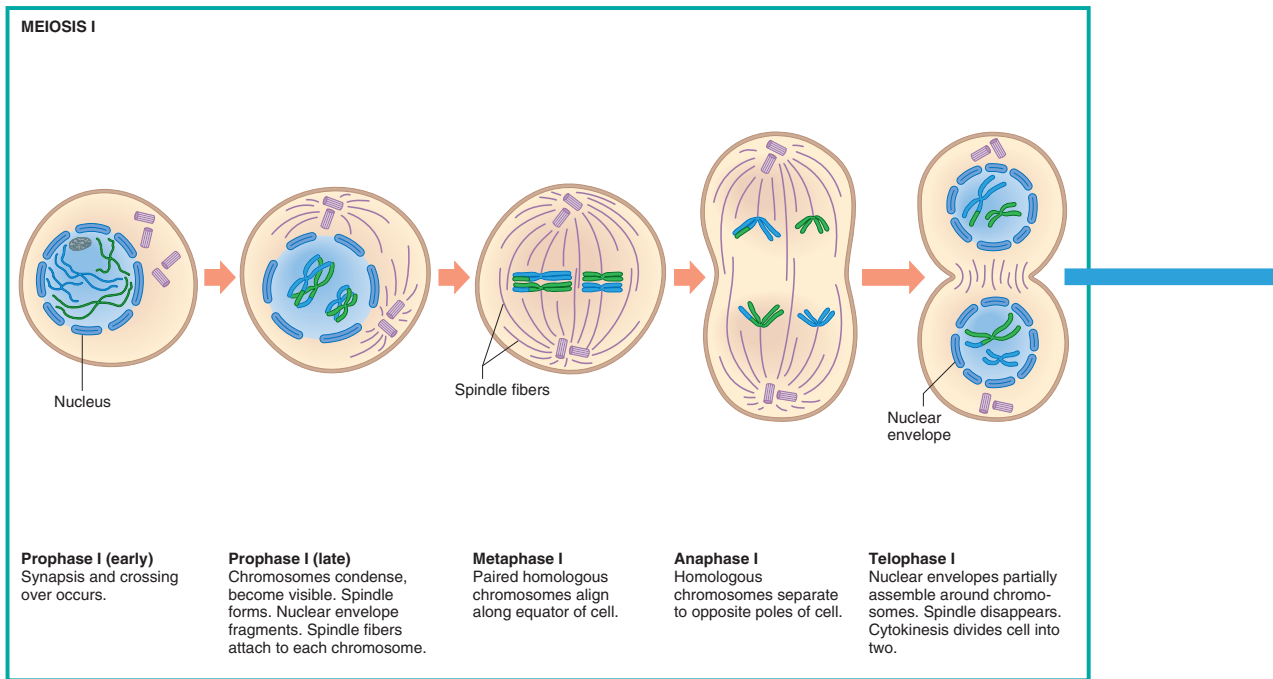


Figure 3.4 Meiosis. An actual human cell undergoing meiosis has 23 chromosome pairs.

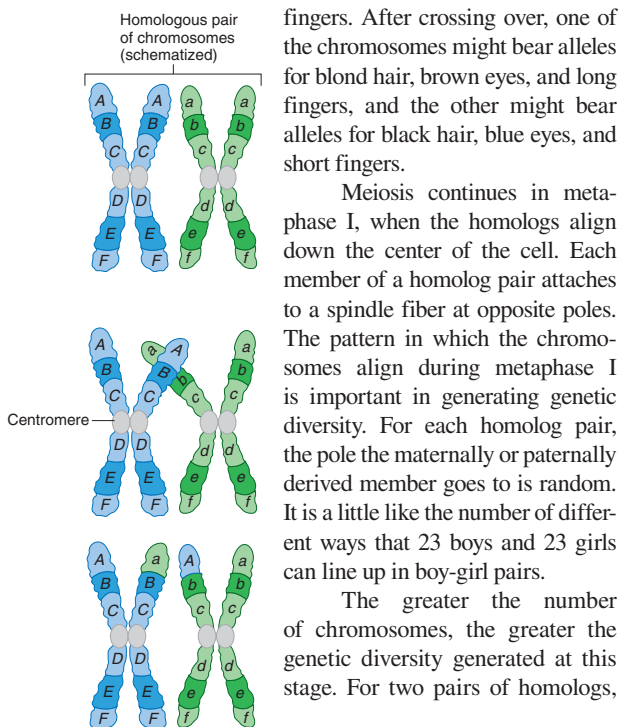


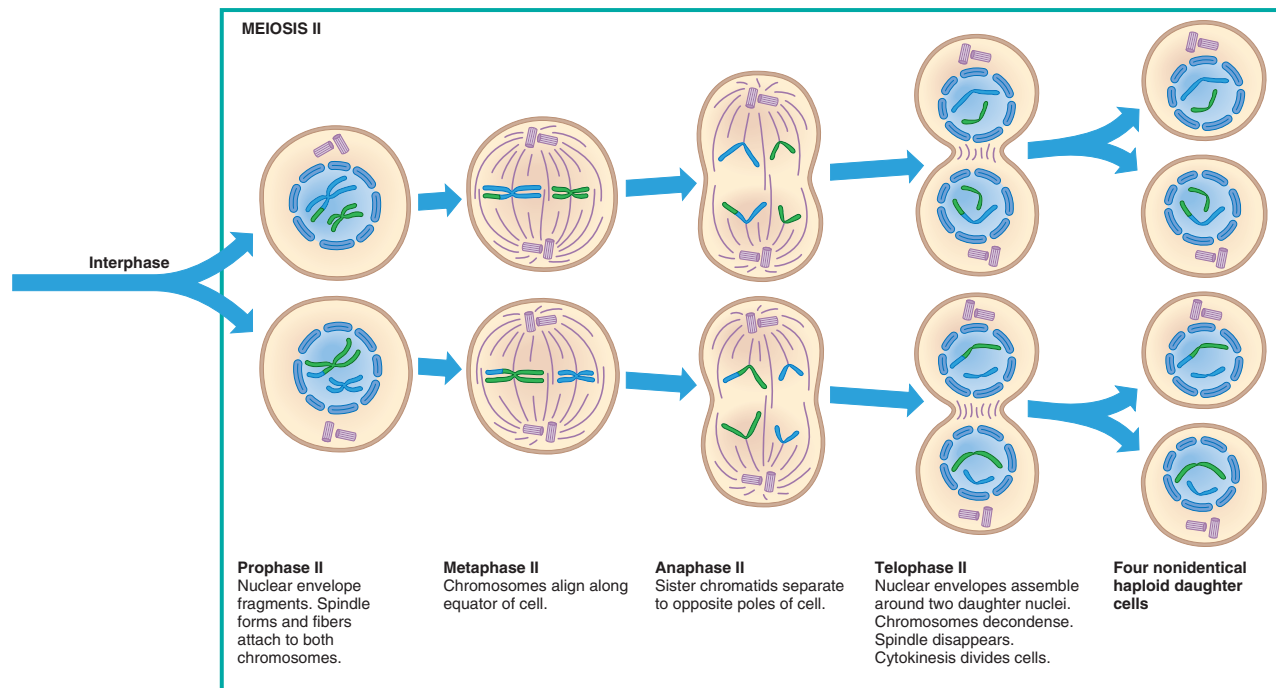
Figure 3.5 Crossing over recombines genes. Crossing over generates genetic diversity by recombining genes and thereby mixing parental traits. The capital and lowercase forms of the same letter represent different variants (alleles) of the same gene. A chromosome actually has hundreds to thousands of genes.

four (2^2) different metaphase alignments are possible. For three pairs of homologs, eight (2^3) different alignments can occur. Our 23 chromosome pairs can line up in 8,388,608 (2^{23}) different ways. This random alignment of chromosomes causes **independent assortment** of the genes that they carry. Independent assortment means that the fate of a gene on one chromosome is not influenced by a gene on a different chromosome (**figure 3.6**). Independent assortment accounts for a basic law of inheritance discussed in chapter 4.

Homologs separate in anaphase I and finish moving to opposite poles by telophase I. These movements establish a haploid set of still-replicated chromosomes at each end of the stretched-out cell. Unlike in mitosis, the centromeres of each homolog in meiosis I remain together. During a second interphase, chromosomes unfold into very thin threads. Proteins are manufactured, but DNA is not replicated a second time. The single DNA replication, followed by the double division of meiosis, halves the chromosome number.

Prophase II marks the start of the second meiotic division. The chromosomes are again condensed and visible. In metaphase II, the replicated chromosomes align down the center of the cell. In anaphase II, the centromeres part, and the newly formed chromosomes, each now in the unreplicated form, move to opposite poles. In telophase II, nuclear envelopes form around the four nuclei, which then separate into individual cells. The net result of meiosis is four haploid cells, each carrying a new assortment of genes and chromosomes that represent a single copy of the genome.

Meiosis generates astounding genetic variety. Any one of a person's more than 8 million possible combinations of



chromosomes can meet with any one of the more than 8 million combinations of a partner, raising potential variability to more than 70 trillion ($8,388,608^2$) genetically unique individuals! Crossing over contributes even more genetic variability.

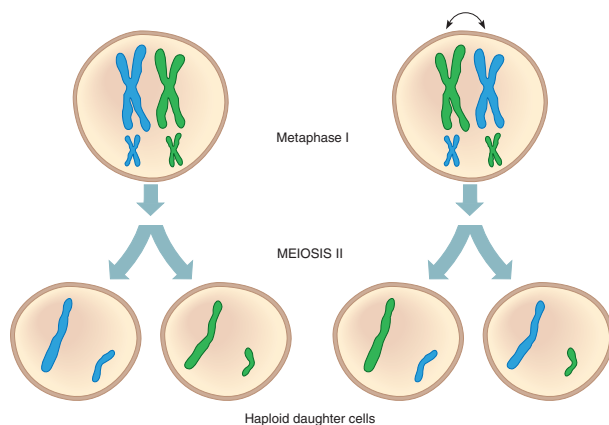


Figure 3.6 Independent assortment. The pattern in which homologs randomly align during metaphase I determines the combination of maternally and paternally derived chromosomes in the daughter cells. Two pairs of chromosomes can align in two different ways to produce four different possibilities in the daughter cells. The potential variability that meiosis generates skyrockets when one considers all 23 chromosome pairs and the effects of crossing over.

Key Concepts

1. The haploid sperm and oocyte are derived from diploid germline cells by meiosis and maturation.
2. Meiosis maintains the chromosome number over generations and mixes gene combinations.
3. In the first meiotic (or reduction) division, the number of replicated chromosomes is halved.
4. In the second meiotic (or equational) division, each of two cells from the first division divides again, yielding four cells from the original one.
5. Chromosome number is halved because the DNA replicates once, but the cell divides twice.
6. Crossing over and independent assortment generate further genotypic diversity by creating new combinations of alleles.

3.3 Gamete Maturation

Meiosis occurs in both sexes, but further steps elaborate the very different-looking sperm and oocyte. Each type of gamete is haploid, but different distributions of other cell components create their distinctions. The cells of the maturing male and female proceed through similar stages, but with sex-specific terminology and different timetables. A male begins manufacturing sperm at puberty and continues throughout life, whereas a female begins meiosis when she is a fetus. Meiosis in the female completes only if a sperm fertilizes an oocyte.

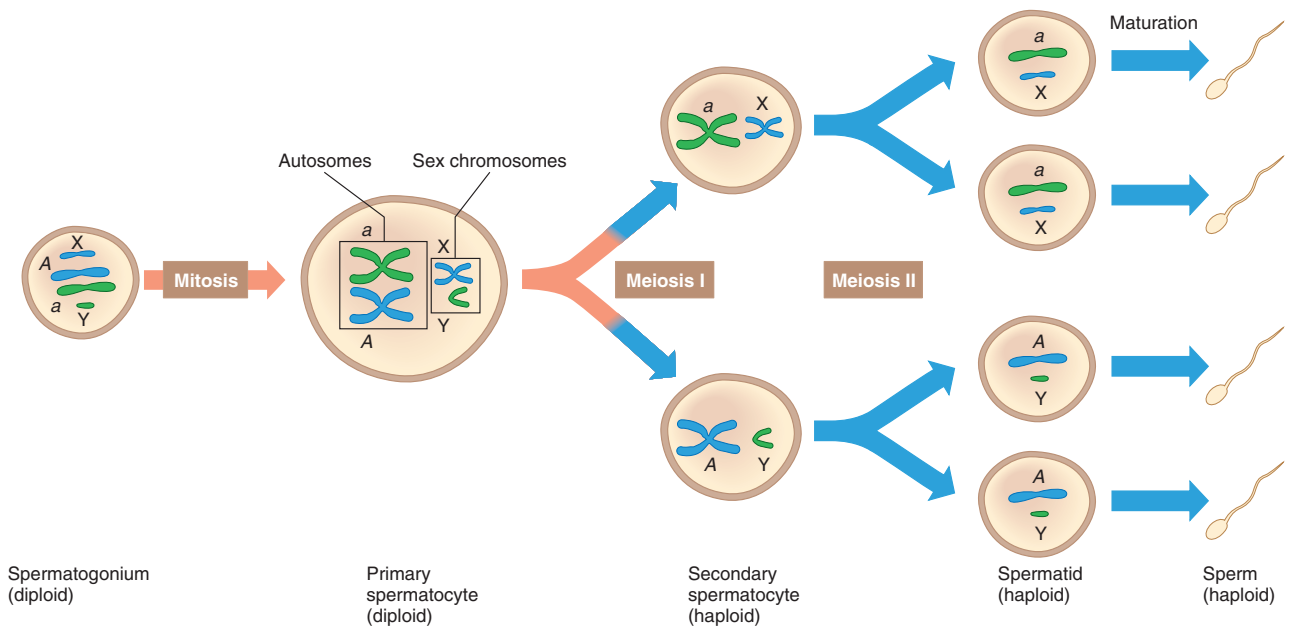


Figure 3.7 Sperm formation (spermatogenesis). Primary spermatocytes have the normal diploid number of 23 chromosome pairs. The large pair of chromosomes represents autosomes (non-sex chromosomes). The X and Y chromosomes are sex chromosomes.

Sperm Formation

Spermatogenesis, the formation of sperm cells, begins in a diploid stem cell called a **spermatogonium** (figure 3.7). This cell divides mitotically, yielding two daughter cells. One continues to specialize into a mature sperm, and the other remains a stem cell, able to self-renew and continually produce more sperm.

Bridges of cytoplasm attach several spermatogonia, and their daughter cells enter meiosis together. As these spermatogonia mature, they accumulate cytoplasm and replicate their DNA, becoming primary spermatocytes.

During reduction division (meiosis I), each primary spermatocyte divides, forming two equal-sized haploid cells called secondary spermatocytes. In meiosis II, each secondary spermatocyte divides to yield two equal-sized spermatids. Each spermatid then develops the characteristic sperm tail, or flagellum. The base of the tail has many mitochondria, which will split ATP molecules to release energy that will propel the sperm inside the female reproductive tract. After spermatid differentiation, some of the cytoplasm connecting the cells falls away, leaving mature, tadpole-shaped spermatozoa (singular, *spermatozoon*), or sperm. **Figure 3.8** presents an anatomical view showing the stages of spermatogenesis within the seminiferous tubules.

A sperm, which is a mere 0.006 centimeter (0.0023 inch) long, must travel about 18 centimeters (7 inches)

to reach an oocyte. Each sperm cell consists of a tail, body or midpiece, and a head region (figure 3.9). A membrane-covered area on the front end, the acrosome, contains enzymes that help the cell penetrate the protective layers around the oocyte. Within the large sperm head, DNA is wrapped around proteins. The sperm's DNA at this time is genetically inactive. A male manufactures trillions of sperm in his lifetime. Although many of these will come close to an oocyte, very few will actually touch one.

Meiosis in the male has built-in protections that help prevent sperm from causing some birth defects. Spermatogonia that are exposed to toxins tend to be so damaged that they never mature into sperm. More mature sperm cells exposed to toxins are often so damaged that they cannot swim.

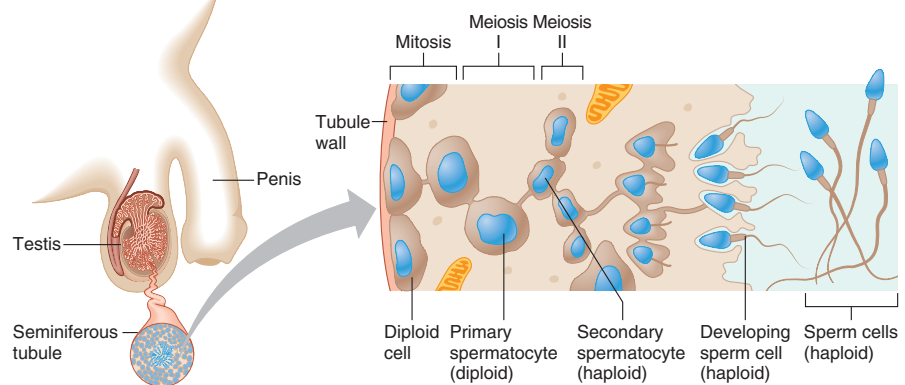


Figure 3.8 Meiosis produces sperm cells. Diploid cells divide through mitosis in the linings of the seminiferous tubules. Some of the daughter cells then undergo meiosis, producing haploid spermatocytes, which differentiate into mature sperm cells.

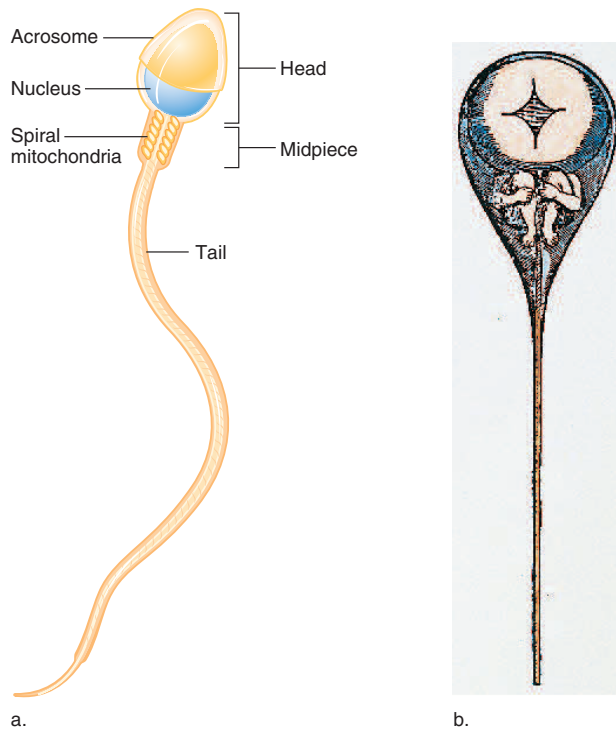


Figure 3.9 Sperm. (a) A sperm has distinct regions that assist in delivering DNA to an oocyte. (b) This 1694 illustration by Dutch histologist Niklass Hartsoeker presents a once-popular hypothesis that a sperm carries a preformed human called a homunculus.

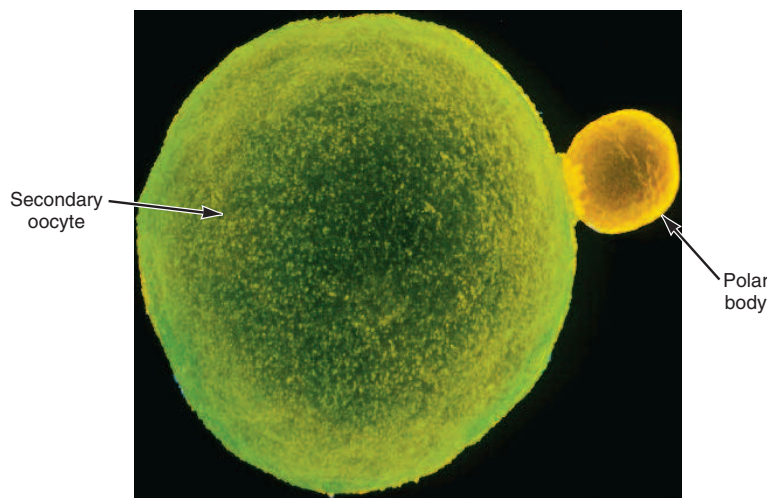


Figure 3.10 Meiosis in a female produces a secondary oocyte and a polar body. Unequal division enables the cell destined to become a fertilized ovum to accumulate most of the cytoplasm and organelles from the primary oocyte, but with only one genome copy. The oocyte accumulates abundant cytoplasm that would have gone into the meiotic product that became the polar body if the division had been equal.

Key Concepts

1. Spermatogonia divide mitotically, yielding one stem cell and one cell that accumulates cytoplasm and becomes a primary spermatocyte.
2. In meiosis I, each primary spermatocyte halves its genetic material to form two secondary spermatocytes.
3. In meiosis II, each secondary spermatocyte divides, yielding two equal-sized spermatids attached by bridges of cytoplasm. Maturing spermatids separate and shed some cytoplasm.
4. A mature sperm has a tail, body, and head, with an enzyme-containing acrosome covering the head.

Oocyte Formation

Meiosis in the female, called **oogenesis** (egg making), begins with a diploid cell, an **oogonium**. Unlike male cells, oogonia are not attached. Instead, follicle cells surround each oogonium. As each oogonium grows, cytoplasm accumulates, DNA replicates, and the cell becomes a primary oocyte. The ensuing meiotic division in oogenesis, unlike the male pathway, produces cells of different sizes.

In meiosis I, the primary oocyte divides into two cells: a small cell with very little cytoplasm, called a first **polar body**, and a much larger cell called a secondary oocyte (**figure 3.10**). Each cell is haploid, with the chromosomes in replicated form. In meiosis II, the tiny first polar body may divide to yield two polar bodies of equal size, with unreplicated chromosomes; or the first polar body may decompose. The secondary oocyte, however, divides unequally in meiosis II to produce another small polar body, with unreplicated chromosomes, and the mature egg cell, or ovum, which contains a large volume of cytoplasm. **Figure 3.11** summarizes meiosis in the female, and **figure 3.12** provides an anatomical view of the process.

Most of the cytoplasm among the four meiotic products in the female ends up in only one cell, the ovum. The woman's body absorbs the polar bodies, which normally play no further role in development. Rarely, a sperm fertilizes a polar body. When this happens, the woman's hormones respond as if she is pregnant, but a disorganized clump of cells that is not an embryo grows for a few weeks, and then leaves the woman's body. This event is a type of miscarriage called a "blighted ovum."

Before birth, a female's million or so oocytes arrest in prophase I. (This means that when your grandmother was pregnant with your mother, the oocyte that would be fertilized and eventually become you was already there.) By puberty, about 400,000 oocytes remain. After puberty, meiosis I continues in one or several oocytes each month, but halts again at metaphase II. In response to specific hormonal cues each month, one ovary releases

a secondary oocyte; this event is ovulation. The oocyte drops into a uterine tube, where waving cilia move it toward the uterus. Along the way, if a sperm penetrates the oocyte membrane, then female meiosis completes, and a fertilized ovum forms. If the secondary oocyte is not fertilized, it degenerates and leaves the body in the menstrual flow, meiosis never completed.

A female ovulates about 400 oocytes between puberty and menopause. However, experiments in mice suggest that stem cells may produce oocytes even past menopause. Most oocytes degrade, because fertilization is so rare. Furthermore, only one in three of the oocytes that do meet and merge with a sperm cell will continue to grow, divide, and specialize to eventually form a new individual.

Key Concepts

1. An oogonium accumulates cytoplasm and replicates its DNA, becoming a primary oocyte.
2. In meiosis I, the primary oocyte divides, forming a small polar body and a large, haploid secondary oocyte.
3. In meiosis II, the secondary oocyte divides, yielding another small polar body and a mature haploid ovum.
4. Oocytes arrest at prophase I until puberty, after which one or several oocytes complete the first meiotic division each month. The second meiotic division completes at fertilization.

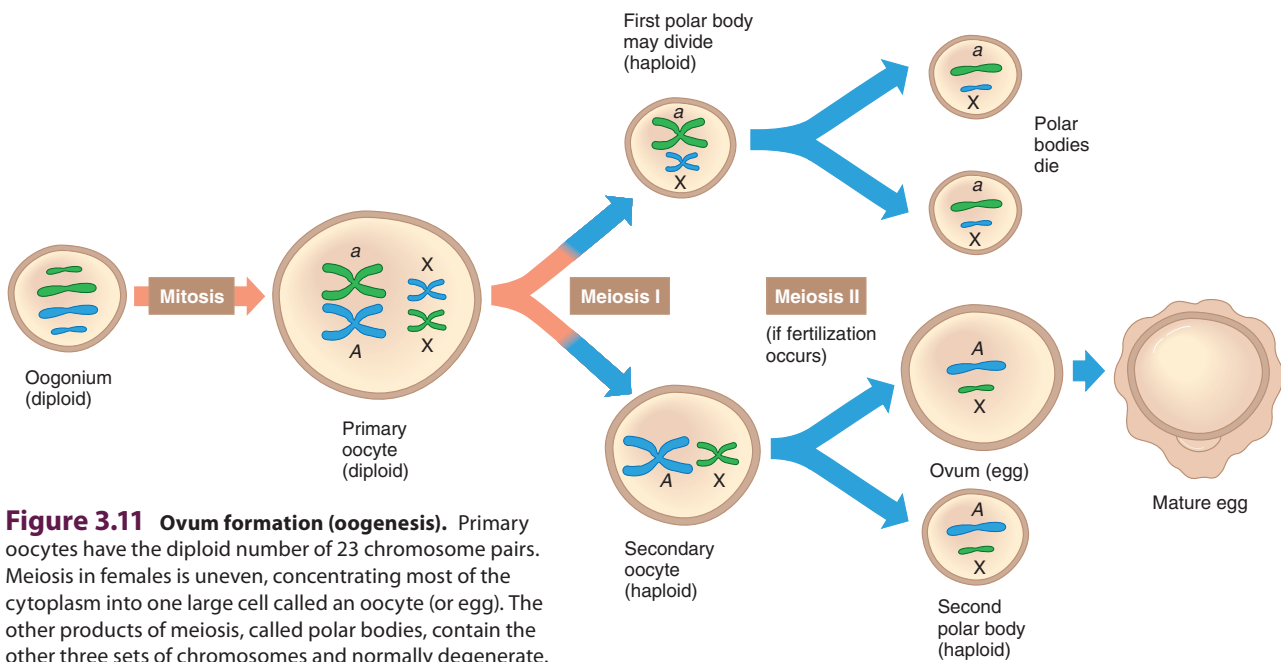


Figure 3.11 Ovum formation (oogenesis). Primary oocytes have the diploid number of 23 chromosome pairs. Meiosis in females is uneven, concentrating most of the cytoplasm into one large cell called an oocyte (or egg). The other products of meiosis, called polar bodies, contain the other three sets of chromosomes and normally degenerate.

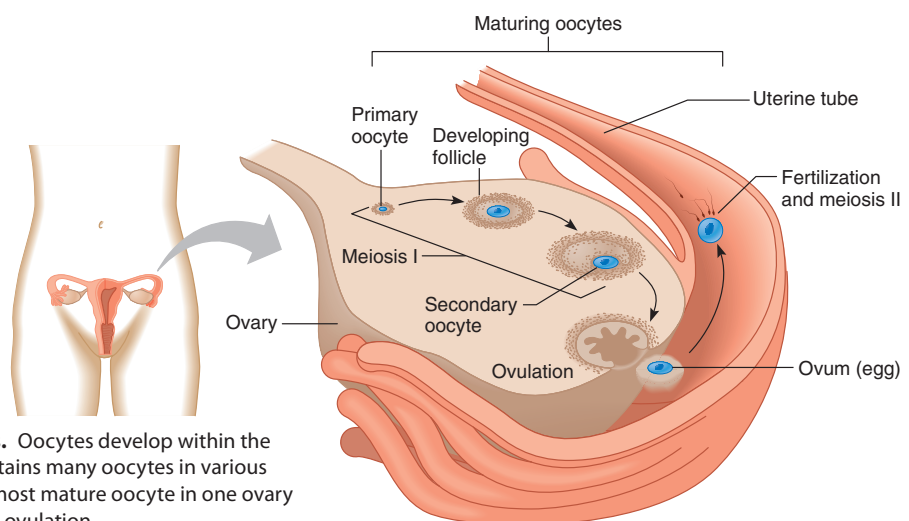


Figure 3.12 The making of oocytes. Oocytes develop within the ovary in protective follicles. An ovary contains many oocytes in various stages of maturation. After puberty, the most mature oocyte in one ovary bursts out each month, in an event called ovulation.

3.4 Prenatal Development

A prenatal human is considered an **embryo** for the first 8 weeks. During this time, rudiments of all body parts form. The embryo in the first week is considered to be in a “preimplantation” stage because it has not yet settled into the uterine lining. Some biologists do not consider a prenatal human an embryo until it begins to develop tissue layers, at about 2 weeks.

Prenatal development after the eighth week is the fetal period, when structures grow and specialize. From the start of the ninth week until birth, the prenatal human organism is a **fetus**.

Fertilization

Hundreds of millions of sperm cells are deposited in the vagina during sexual intercourse. A sperm cell can survive in the woman’s body for up to 3 days, but the oocyte can only be fertilized in the 12 to 24 hours after ovulation.

The woman’s body helps sperm reach an oocyte. A process in the female called capacitation chemically activates sperm, and the oocyte secretes a chemical that attracts sperm. Sperm are also assisted by contractions of the female’s muscles and by their moving tails. Still, only 200 or so sperm come near the oocyte.

A sperm first contacts a covering of follicle cells, called the corona radiata, that guards a secondary oocyte. The sperm’s acrosome then bursts, releasing enzymes that bore through a protective layer of glycoprotein (the zona pellucida) beneath the corona radiata. Fertilization, or conception, begins when the outer membranes of the sperm and secondary

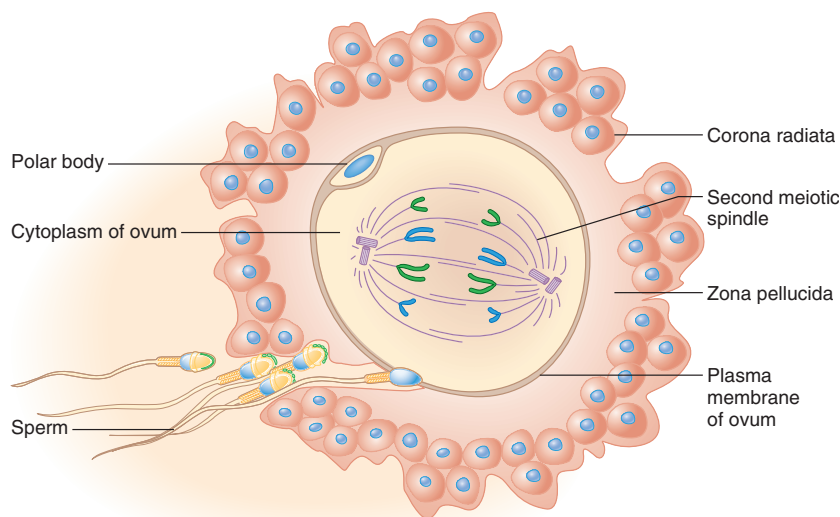
oocyte meet (**figure 3.13**). The encounter is dramatic. A wave of electricity spreads physical and chemical changes across the entire oocyte surface—changes that keep other sperm out. More than one sperm can enter an oocyte, but the resulting cell has too much genetic material for development to follow.

Usually only the sperm’s head enters the oocyte. Within 12 hours of the sperm’s penetration, the ovum’s nuclear membrane disappears, and the two sets of chromosomes, called pronuclei, approach one another. Within each pronucleus, DNA replicates. Fertilization completes when the two genetic packages meet and merge, forming the genetic instructions for a new individual. The fertilized ovum is called a **zygote**. The *Bioethics: Choices for the Future* reading describes cloning, which is a way to start development without a fertilized egg.

Cleavage and Implantation

About a day after fertilization, the zygote divides by mitosis, beginning a period of frequent cell division called **cleavage** (**figure 3.14**). The resulting early cells are called **blastomeres**. When the blastomeres form a solid ball of sixteen or more cells, the embryo is called a **morula** (Latin for “mulberry,” which it resembles).

During cleavage, organelles and molecules from the secondary oocyte’s cytoplasm still control cellular activities, but some of the embryo’s genes begin to function. The ball of cells hollows out, and its center fills with fluid, creating a **blastocyst**—the “cyst” refers to the fluid-filled center. Some of the cells form a clump on the inside lining called the **inner cell mass**. Formation of the inner cell mass is the first event



a.



b.

Figure 3.13 Fertilization. (a) Fertilization by a sperm cell induces the oocyte (arrested in metaphase II) to complete meiosis. Before fertilization occurs, the sperm’s acrosome bursts, spilling enzymes that help the sperm’s nucleus enter the oocyte. (b) Only about 200 sperm come near an oocyte, and only one can enter.

Bioethics: Choices for the Future

Why a Clone Is Not an Exact Duplicate

Cloning is the creation of a genetic replica of an individual. This is in contrast to normal reproduction and development, in which genetic material from two individuals combines. In fiction, scientists have cloned Nazis, politicians, dinosaurs, and organ donors for wealthy people. Real scientists have cloned sheep, mice, cats, pigs, dogs, and amphibians.

Cloning transfers a nucleus from a somatic cell into an oocyte whose nucleus has been removed, which then develops. The technique is more accurately called “somatic cell nuclear transfer” (SCNT) or just “nuclear transfer.” However, nuclear transfer cannot produce an exact replica of a person, for several reasons:

- **Premature cellular aging.** In some species, telomeres of chromosomes in the donor nucleus are shorter than those in the recipient cell (see chapter 2). Premature aging, seen in shortened telomeres, may be why the first cloned mammal, Dolly, died young of a severe respiratory infection.
- **Altered gene expression.** In normal development, for some genes, one copy is turned off, depending upon which parent transmits it. That is, some genes must be inherited from either the father or the mother to act. This phenomenon is called genomic imprinting. In cloning, genes in a donor nucleus skip passing through a sperm or oocyte, and thus are not imprinted. Lack of imprinting may cause cloned animals to be unusually large. Regulation of gene expression is abnormal at many times during prenatal development (see chapter 5).
- **More mutations.** DNA from a donor cell has had years to accumulate mutations. Such a mutation might not be noticeable in one of millions of somatic cells, but it could be devastating if that nucleus is used to program development of a new individual (see chapter 11).
- **X inactivation.** At a certain time in early prenatal development in female mammals, one X chromosome is inactivated. Whether that X chromosome is from the mother or the father occurs

at random in each cell, creating an overall mosaic pattern of expression for genes on the X chromosome. The pattern of X inactivation of a female clone would probably not match that of her nucleus donor, because X inactivation occurs in the embryo, not the first cell (see chapter 6).

- **Mitochondrial DNA.** Mitochondria contain DNA. A clone’s mitochondria descend from the recipient oocyte, not from the donor cell, because they are in the cytoplasm, not the nucleus.

The environment is another powerful factor in why a clone isn’t an identical copy. For example, coat color patterns differ in cloned calves and cats. When the animals were embryos, cells destined to produce pigment moved in a unique way in each individual, producing different color patterns. In humans, experience, nutrition, stress, exposure to infectious disease, and many other factors join our genes in molding who we are.



Cloned cats. A company called “Genetic Savings and Clone” tried to sell cloned cats for \$50,000, but lowered the price to only \$32,000 when customers were scarce. The company went out of business in 2006.

that distinguishes cells from each other in terms of their relative positions, other than the inside and outside of the morula. The cells of the inner cell mass will continue developing to form the embryo. (Cells that can be used to generate embryonic stem cells come from the 8-celled cleavage embryo or the inner cell mass of a 5-day blastocyst, shown in figure 2.25.)

A week after conception, the blastocyst begins to nestle into the woman’s uterine lining (endometrium). This event, called implantation, takes about a week. As it starts, the outermost cells of the blastocyst, called the trophoblast, secrete the “pregnancy hormone” human chorionic gonadotropin (hCG), which prevents menstruation. hCG detected in a woman’s urine or blood is one sign of pregnancy.

Key Concepts

1. Following sexual intercourse, sperm are capacitated and drawn to the secondary oocyte.
2. Acrosomal enzymes assist the sperm’s penetration of the oocyte. Chemical and electrical changes in the oocyte’s surface block additional sperm.
3. The two sets of chromosomes meet, forming a zygote.
4. Cleavage cell divisions form a morula and then a blastocyst.
5. The outer layer of cells invades and implants in the uterine lining.
6. The inner cell mass develops into the embryo.
7. Certain blastocyst cells secrete hCG.

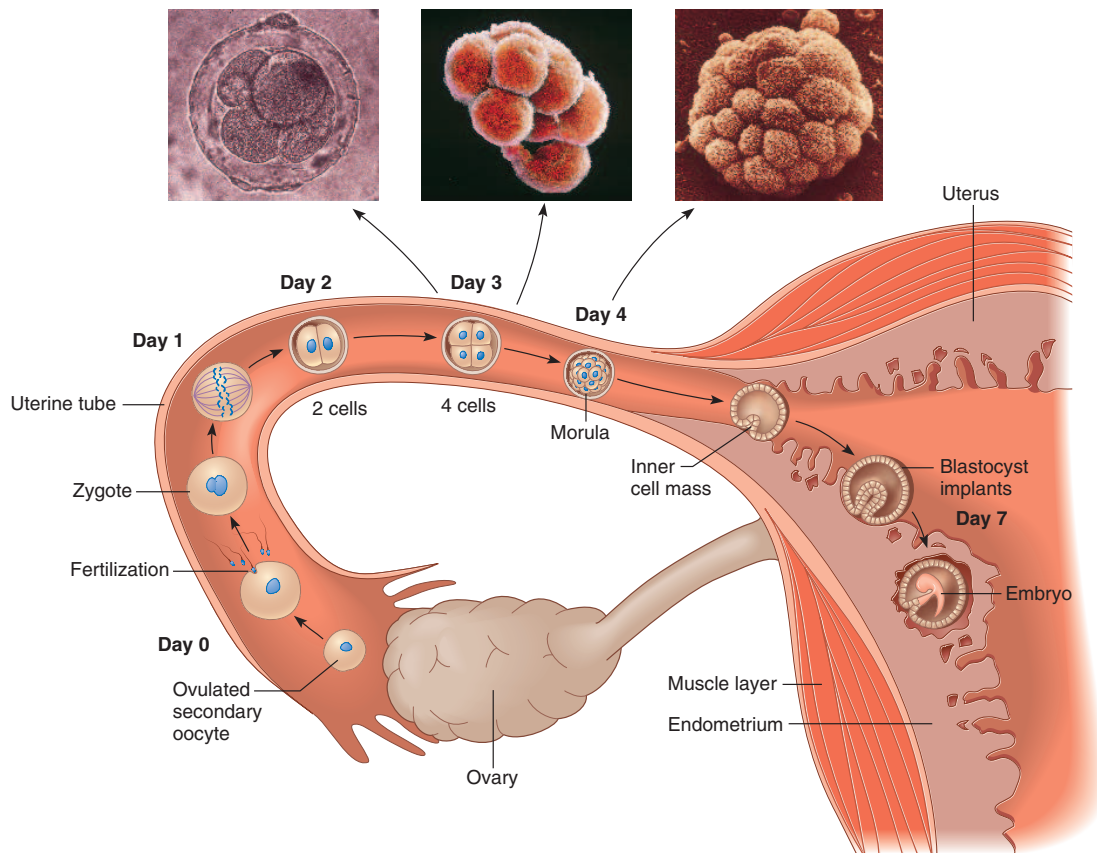


Figure 3.14 Cleavage: From ovulation to implantation. The zygote forms in the uterine tube when a sperm nucleus fuses with the nucleus of an oocyte. The first divisions proceed while the zygote moves toward the uterus. By day 7, the zygote, now called a blastocyst, begins to implant in the uterine lining.

The Embryo Forms

During the second week of prenatal development, a space called the amniotic cavity forms between the inner cell mass and the outer cells anchored to the uterine lining. Then the inner cell mass flattens into a two-layered embryonic disc. The layer nearest the amniotic cavity is the **ectoderm**; the inner layer, closer to the blastocyst cavity, is the **endoderm**. Shortly after, a third layer, the **mesoderm**, forms in the middle. This three-layered structure is called the primordial embryo, or the **gastrula** (figure 3.15).

Once these three layers, called **primary germ layers**, form, many cells become “determined” or fated, to develop as a specific cell type. Each primary germ layer gives rise to certain structures. Cells in the ectoderm become skin, nervous tissue, or parts of certain glands. Endoderm cells form parts of the liver and pancreas and the linings of many organs. The middle layer of the embryo, the mesoderm, forms many structures, including muscle, connective tissues, the reproductive organs, and the kidneys.

Genes called homeotics control how the embryo develops parts in the right places. Mutations in these genes cause some

very interesting conditions, including forms of mental retardation, autism, and blindness. Figure 16.1b shows one that disrupts hand development. The homeotic mutations were originally studied in fruit flies that had legs growing where their antennae should be. The author did her graduate work on these flies, never suspecting that the mutations had counterparts in humans.

Table 3.2 summarizes the stages of early human prenatal development.

Supportive Structures Form

As an embryo develops, structures form that support and protect it. These include chorionic villi, the placenta, the yolk sac, the allantois, the umbilical cord, and the amniotic sac.

By the third week after conception, finger-like outgrowths called chorionic villi extend from the area of the embryonic disc close to the uterine wall, and these project into pools of the woman’s blood. Her blood system and the embryo’s are separate, but nutrients and oxygen diffuse across the chorionic villi from her circulation to the embryo, and wastes leave the embryo’s circulation and enter the woman’s circulation to be excreted.

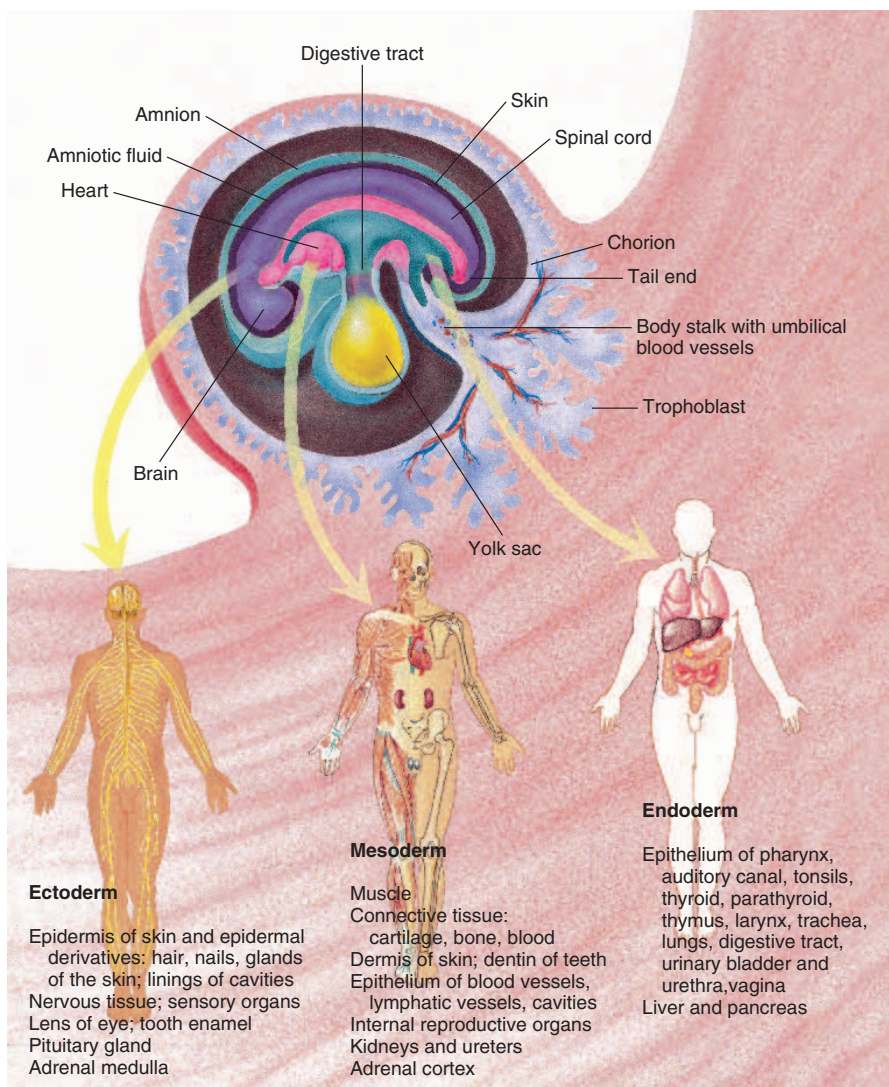


Figure 3.15 The primordial embryo. When the three primary germ layers of the embryo form at gastrulation, many cells become “fated” to follow a specific developmental pathway. Each layer retains stem cells as the organism develops. Under certain conditions, these cells may produce daughter cells that can specialize as many cell types.

Table 3.2 Stages and Events of Early Human Prenatal Development

Stage	Time Period	Principal Events
Fertilized ovum	12–24 hours following ovulation	Oocyte fertilized; zygote has 23 pairs of chromosomes and is genetically distinct
Cleavage	30 hours to third day	Mitosis increases cell number
Morula	Third to fourth day	Solid ball of cells
Blastocyst	Fifth day through second week	Hollowed ball forms trophoblast (outside) and inner cell mass, which implants and flattens to form embryonic disc
Gastrula	End of second week	Primary germ layers form

By 10 weeks, the placenta is fully formed. The placenta is an organ that links woman and fetus for the rest of the pregnancy. The placenta secretes hormones that maintain pregnancy and alter the woman’s metabolism to send nutrients to the fetus.

Other structures nurture the developing embryo. The yolk sac manufactures blood cells, as does the allantois, a membrane surrounding the embryo that gives rise to the umbilical blood vessels. The umbilical cord forms around these vessels and attaches to the center of the placenta. Toward the end of the embryonic period, the yolk sac shrinks, and the amniotic sac swells with fluid that cushions the embryo and maintains a constant temperature and pressure. The amniotic fluid contains fetal urine and cells.

Two of the supportive structures that develop during pregnancy provide the material for prenatal tests (see figure 13.5), discussed in chapter 13. Chorionic villus sampling examines chromosomes from cells snipped off the chorionic villi at 10 weeks. Because the villi cells and the embryo’s cells come from the same fertilized ovum, an abnormal chromosome in villi cells should also be in the embryo. In amniocentesis, a sample of amniotic fluid is taken and fetal cells in it are examined for biochemical, genetic, and chromosomal anomalies.

The umbilical cord is another prenatal structure that has medical applications. In addition to the cord blood banks mentioned in *Bioethics: Choices for the Future* in chapter 2, the cells of the cord itself are valuable. When cultured in the same “cocktails” used for embryonic stem cells, cord cells can differentiate as cells from any of the three primary germ layers, including bone, fat, nerve, cartilage, and muscle cells. Stem cells from the cord are used to treat a respiratory disease of newborns that scars and inflames the lungs. The stem cells become two types of needed lung cells: the type that secretes surfactant, which is the chemical that inflates the microscopic air sacs, and the type that

exchanges oxygen for carbon dioxide. Stem cells from umbilical cords are abundant, easy to obtain and manipulate, and can become almost any cell type.

Key Concepts

1. Primary germ layers form in the second week. Cells in a specific germ layer later become parts of particular organ systems as a result of differential gene expression.
2. During week 3, chorionic villi extend toward the maternal circulation, and the placenta begins to form.
3. Nutrients and oxygen enter the embryo, and wastes pass from the embryo into the maternal circulation.
4. The yolk sac and allantois manufacture blood cells, the umbilical cord forms, and the amniotic sac expands with fluid.

Multiples

Twins and other multiples arise early in development. Twins are either fraternal or identical. Fraternal, or **dizygotic** (DZ), twins result when two sperm fertilize two oocytes. This can happen if ovulation occurs in two ovaries in the same month, or if two oocytes leave the same ovary and are both fertilized. DZ twins are no more alike than any two siblings, although they share a very early environment in the uterus. The tendency to have DZ twins may run in families if the women sometimes ovulate two oocytes in a month.

Identical, or **monozygotic** (MZ), twins descend from a single fertilized ovum and therefore are genetically identical. They are natural clones. Three types of MZ twins can form, depending upon when the fertilized ovum or very early embryo splits (**figure 3.16**). This difference in timing determines which supportive structures the twins share. About a third of all MZ twins have completely separate chorions and amnions, and about two-thirds share a chorion but have separate amnions. Slightly fewer than 1 percent of MZ twins share both amnion and chorion. (The amnion is the sac that contains fluid that surrounds the fetus. The chorion develops into the placenta.) These differences may expose the different types of MZ twins to slightly different uterine environments. For example, if one chorion develops more attachment sites to the maternal circulation, one twin may receive more nutrients and gain more weight than the other.

In 1 in 50,000 to 100,000 pregnancies, an embryo divides into twins after the point at which the two groups of cells can develop as two individuals, between days 13 and 15. The result is conjoined or “Siamese” twins. The latter name comes from Chang and Eng Bunker, who were born in Thailand, then called Siam, in 1811. They were joined by a band of tissue from the navel to the breastbone, and could easily have been separated today. Chang and Eng lived for 63 years, attached. They fathered 22 children and divided each week between their wives.

For Abigail and Brittany Hensel, shown in **figure 3.17**, the separation occurred after day 9 of development, but before

day 14. Biologists know this because the girls’ shared organs have derivatives of ectoderm, mesoderm, and endoderm; that is, when the lump of cells divided incompletely, the three primary germ layers had not yet completely sorted themselves out. The Hensel girls are extremely rare “incomplete twins.” They are “dicephalic,” which means that they have two heads. They are very much individuals.

Each girl has her own neck, head, heart, stomach, gallbladder, and lungs. Each has one leg and one arm, and a third arm between their heads was surgically removed. Each girl also has her own nervous system! The twins share a large liver, a single bloodstream, and all organs below the navel. They have three kidneys. Because at birth Abby and Brittany were strong and healthy, doctors suggested surgery to separate them. But their parents, aware from other cases that only one child would likely survive a separation, chose to let their daughters be.

As teens, Abby and Brittany are glad their parents did not choose to separate them, because they would have been unable to walk or run, as they can today. They enjoy kickball, volleyball, basketball, and cycling. Like any teen girls, they have distinctive tastes in clothing and in food.

MZ twins occur in 3 to 4 pregnancies per 1,000 births worldwide. In North America, twins occur in about 1 in 81 pregnancies, which means that 1 in 40 of us is a twin. However, not all twins survive to be born. One study of twins detected early in pregnancy showed that up to 70 percent of the eventual births are of a single child. This is called the “vanishing twin” phenomenon.

Key Concepts

1. Dizygotic twins arise from two fertilized ova.
2. Monozygotic twins arise from a single fertilized ovum and may share supportive structures.

The Embryo Develops

As the days and weeks of prenatal development proceed, different rates of cell division in different parts of the embryo fold the forming tissues into intricate patterns. In a process called embryonic induction, the specialization of one group of cells causes adjacent groups of cells to specialize. Gradually, these changes mold the three primary germ layers into organs and organ systems. Organogenesis is the transformation of the simple three layers of the embryo into distinct organs. During the weeks of organogenesis, the developing embryo is particularly sensitive to environmental influences such as chemicals and viruses.

During the third week of prenatal development, a band called the primitive streak appears along the back of the embryo. The primitive streak gradually elongates to form an axis that other structures organize around as they develop. The primitive streak eventually gives rise to connective tissue progenitor cells and the notochord, which is a structure that forms the basic framework of the skeleton. The notochord induces a sheet of

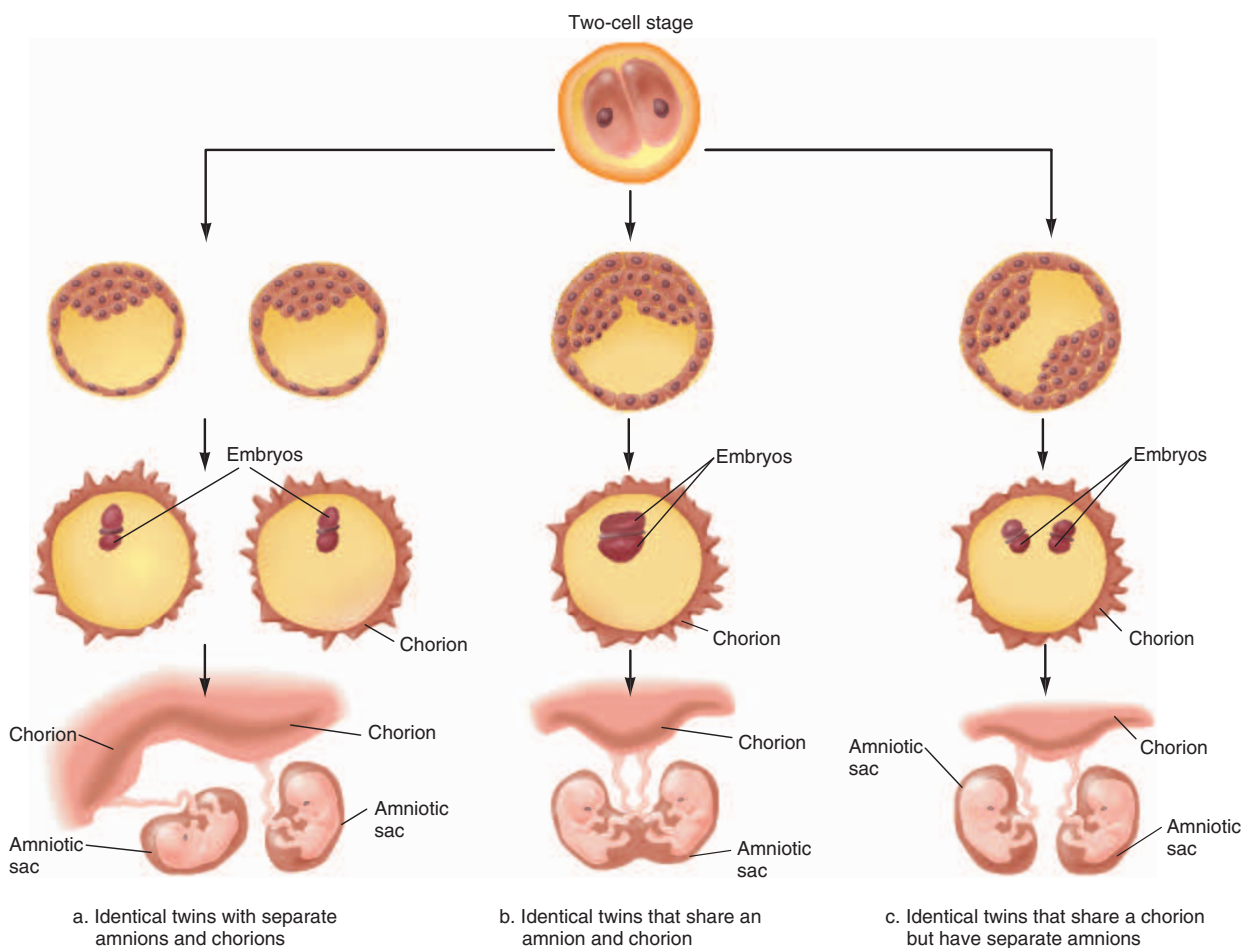


Figure 3.16 Types of identical twins. Identical twins originate at three points in development. **(a)** In about one-third of identical twins, separation of cells into two groups occurs before the trophoblast forms on day 5. These twins have separate chorions and amnions. **(b)** About 1 percent of identical twins share a single amnion and chorion, because the tissue splits into two groups after these structures have already formed. **(c)** In about two-thirds of identical twins, the split occurs after day 5 but before day 9. These twins share a chorion but have separate amnions. Fraternal twins result from two sperm fertilizing two secondary oocytes. These twins develop their own amniotic sacs, yolk sacs, allantois, placentae, and umbilical cords.

overlying ectoderm to fold into the hollow **neural tube**, which develops into the brain and spinal cord (central nervous system). If the neural tube does not completely zip up by day 20, a birth defect called a neural tube defect (NTD) occurs. As a result, parts of the brain or spinal cord protrude from the open head or spine, and body parts below the defect are not innervated. The person is paralyzed from the point of the NTD down. Surgery can correct some NTDs (see *Bioethics: Choices for the Future* Box in chapter 15). Lack of the B vitamin folic acid can cause NTDs in embryos with a genetic susceptibility. For this reason, the U.S. government adds the vitamin to grains, and pregnant women take supplements. A blood test during the 15th week of pregnancy detects a substance from the fetus's liver called alpha fetoprotein (AFP) that leaks at an abnormally rapid rate into the woman's circulation if there is an NTD.

Some nations designate day 14 of prenatal development and primitive streak formation as the point beyond which they

ban research on the human embryo. The reasoning is that the primitive streak is the first sign of a nervous system and day 14 is when implantation completes.

Appearance of the neural tube marks the beginning of organ development. Shortly after, a reddish bulge containing the heart appears. The heart begins to beat around day 18, and this is easily detectable by day 22. Soon the central nervous system starts to form.

The fourth week of embryonic existence is one of spectacularly rapid growth and differentiation (**figure 3.18**). Arms and legs begin to extend from small buds on the torso. Blood cells form and fill primitive blood vessels. Immature lungs and kidneys begin to develop.

By the fifth and sixth weeks, the embryo's head appears to be too large for the rest of its body. Limbs end in platelike structures with tiny ridges, and gradually apoptosis sculpts the fingers and toes. The eyes are open, but they do not yet have



Figure 3.17 Conjoined twins. Abby and Brittany Hensel are the result of incomplete twinning during the first 2 weeks of prenatal development.

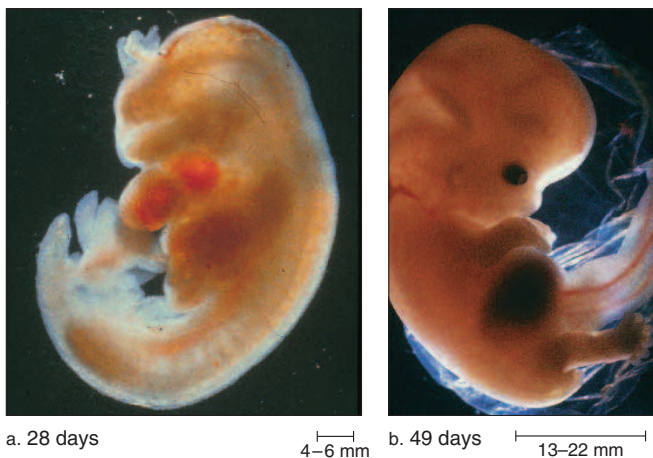


Figure 3.18 Human embryos at (a) 28 days, and (b) 49 days.

lids or irises. By the seventh and eighth weeks, a skeleton composed of cartilage forms. The embryo is now about the length and weight of a paper clip. At eight weeks of gestation, the prenatal human has rudiments of all of the structures that will be present at birth. It is now a fetus.

Key Concepts

1. During week 3, the primitive streak appears, followed rapidly by the notochord, neural tube, heart, central nervous system, limbs, digits, facial features, and other organ rudiments.
2. By week 8, all of the organs that will be present in the newborn have begun to develop.

The Fetus Grows

During the fetal period, body proportions approach those of a newborn. Initially, the ears lie low, and the eyes are widely spaced. Bone begins to replace the softer cartilage. As nerve and muscle functions become coordinated, the fetus moves.

Sex is determined at conception, when a sperm bearing an X or Y chromosome meets an oocyte, which always carries an X chromosome. An individual with two X chromosomes is a female, and one with an X and a Y is a male. A gene on the Y chromosome, called *SRY* (for “sex-determining region of the Y”), determines maleness.

Differences between the sexes do not appear until week 6, after the *SRY* gene is expressed in males. Male hormones then stimulate male reproductive organs and glands to differentiate from existing, indifferent structures. In a female, the indifferent structures of the early embryo develop as female organs and glands, under the control of other genes. Differences may be noticeable on ultrasound scans by 12 to 15 weeks. Sexual development is discussed further in chapter 6.

By week 12, the fetus sucks its thumb, kicks, makes fists and faces, and has the beginnings of teeth. It breathes amniotic fluid in and out, and urinates and defecates into it. The first trimester (three months) of pregnancy ends.

By the fourth month, the fetus has hair, eyebrows, lashes, nipples, and nails. By 18 weeks, the vocal cords have formed, but the fetus makes no sound because it doesn’t breathe air. By the end of the fifth month, the fetus curls into a head-to-knees position. It weighs about 454 grams (1 pound). During the sixth month, the skin appears wrinkled because there isn’t much fat beneath it, and turns pink as capillaries fill with blood. By the end of the second trimester, the woman feels distinct kicks and jabs and may even detect a fetal hiccup. The fetus is now about 23 centimeters (9 inches) long.

In the final trimester, fetal brain cells rapidly link into networks as organs elaborate and grow. A layer of fat forms beneath the skin. The digestive and respiratory systems mature last, which is why infants born prematurely often have difficulty digesting milk and breathing. Approximately 266 days after a single sperm burrowed its way into an oocyte, a baby is ready to be born.

The birth of a healthy baby is against the odds. Of every 100 secondary oocytes exposed to sperm, 84 are fertilized. Of these 84, 69 implant in the uterus, 42 survive one week or longer, 37 survive 6 weeks or longer, and only 31 are born alive. Of the fertilized ova that do not survive, about half have chromosomal abnormalities that cause problems too severe for development to proceed.

Key Concepts

1. During the fetal period, structures grow, specialize, and begin to interact.
2. Bone replaces cartilage in the skeleton, body growth catches up with the head, and sex organs become more distinct.
3. In the final trimester, the fetus moves and grows rapidly, and fat fills out the skin.

3.5 Birth Defects

Certain genetic abnormalities or toxic exposures can affect development in an embryo or fetus, causing birth defects. Only a genetic birth defect can be passed to future generations. Although development can be derailed in many ways, about 97 percent of newborns appear healthy at birth.

The Critical Period

The specific nature of a birth defect reflects the structures developing when the damage occurs. The time when genetic abnormalities, toxic substances, or viruses can alter a specific structure is its **critical period** (figure 3.19). Some body parts, such as fingers and toes, are sensitive for short periods of time. In contrast, the brain is sensitive throughout prenatal development, and connections between nerve cells continue to change throughout life. Because of the brain's continuous critical period, many birth defect syndromes include learning disabilities or mental retardation.

About two-thirds of all birth defects arise from a disruption during the embryonic period. More subtle defects, such as learning disabilities, that become noticeable only after infancy are often caused by interventions during the fetal period. A disruption in the first trimester might cause mental retardation; in the seventh month of pregnancy, it might cause difficulty in learning to read.

Some birth defects can be attributed to an abnormal gene that acts at a specific point in prenatal development. In a rare inherited condition called phocomelia (MIM 276826), for example, a mutation halts limb development from the third to the fifth week of the embryonic period, causing “flippers” to develop in place of arms and legs. The risk that a genetically caused birth defect will affect a particular family member can be calculated.

Many birth defects are caused not by mutant genes but by toxic substances the pregnant woman encounters. These

environmentally caused problems will not affect other family members unless they, too, are exposed to the environmental trigger. Chemicals or other agents that cause birth defects are called **teratogens** (Greek for “monster-causing”). While it is best to avoid teratogens while pregnant, some women may need to continue to take potentially teratogenic drugs to maintain their own health.

Teratogens

Most drugs are not teratogens. Whether or not exposure to a particular drug causes birth defects may depend upon a woman's genes. For example, certain variants of a gene that control the body's use of an amino acid called homocysteine affect whether or not the medication valproic acid causes birth defects. Valproic acid is used to prevent seizures and symptoms of bipolar disorder. Rarely, it can cause NTDs, heart defects, hernias, and club foot. Women can be tested for this gene variant (*MTHFR C677T*, MIM 607093) and if they have it, switch to a different medication when they try to conceive. Experiments using stem cells are investigating exactly how valproic acid disrupts development.

Thalidomide

The idea that the placenta protects the embryo and fetus against harmful substances was tragically disproven between 1957 and 1961, when 10,000 children were born in Europe with what seemed, at first, to be phocomelia. Doctors realized that this genetic disorder is very rare, and therefore couldn't be the cause of the sudden problem. They discovered that the mothers had all taken a mild tranquilizer, thalidomide, to alleviate nausea

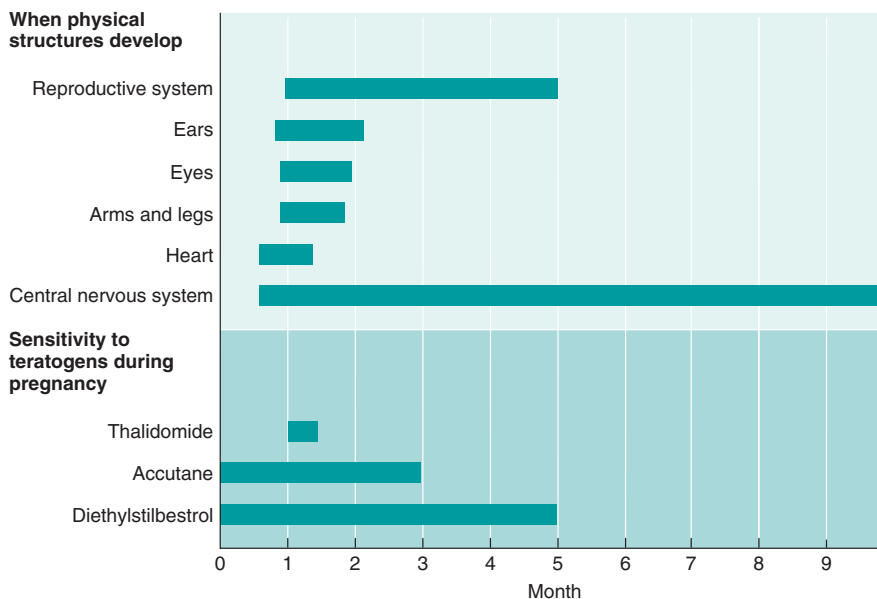


Figure 3.19 Critical periods of development. The nature of a birth defect resulting from drug exposure depends upon which structures were developing at the time of exposure. The time when a particular structure is vulnerable is called its critical period. Accutane is an acne medication that causes cleft palate and eye, brain, and heart defects. Diethylstilbestrol (DES) was used in the 1950s to prevent miscarriage. It caused vaginal cancer in some “DES daughters.” Thalidomide was used to prevent morning sickness.

early in pregnancy, during the critical period for limb formation. Many “thalidomide babies” were born with incomplete or missing legs and arms.

The United States was spared from the thalidomide disaster because an astute government physician noted the drug’s adverse effects on laboratory monkeys. Still, several “thalidomide babies” were born in South America in 1994, where pregnant women were given the drug. In spite of its teratogenic effects, thalidomide is used to treat leprosy, AIDS, and certain blood and bone marrow cancers.

Cocaine

When cocaine use soared in the late 1980s and 1990s, physicians predicted a wave of birth defects in children who were affected by their mothers’ cocaine use while pregnant. However, analysis of studies on more than 4,000 of these children indicate that effects, if any, are mild, and are less severe than those of prenatal exposure to alcohol or tobacco. Cocaine may slow fetal growth, particularly of the head, but the babies quickly catch up. No effects were seen on either language skills or IQ. Children exposed to cocaine in the uterus may have difficulty concentrating and behave poorly, but it is difficult to tease out whether this is due to the past drug exposure, or to the problems associated with poverty, such as malnutrition and hunger. Researchers are continuing to follow the progress of these young people to see if problems develop that can be linked to prenatal cocaine exposure.

Cigarettes

Chemicals in cigarette smoke stress a fetus. Carbon monoxide crosses the placenta and prevents the fetus’s hemoglobin molecules from adequately binding oxygen. Other chemicals in smoke block nutrients. Smoke-exposed placentas lack important growth factors, causing poor growth before and after birth. Cigarette smoking during pregnancy increases the risk of spontaneous abortion, stillbirth, prematurity, and low birth weight.

Alcohol

A pregnant woman who has just one or two alcoholic drinks a day, or perhaps a large amount at a single crucial time, risks fetal alcohol syndrome (FAS) in her unborn child. Tests for gene variants that encode proteins that regulate alcohol metabolism may be able to predict which women and fetuses are at elevated risk for developing FAS, but until these tests are validated, pregnant women are advised to avoid all alcohol.

A child with FAS has a characteristic small head and a flat face (**figure 3.20**). Growth is slow before and after birth. Teens and young adults who have FAS are short and have small heads. More than 80 percent of them retain the facial characteristics of a young child with FAS.

The long-term mental effects of prenatal alcohol exposure are more severe than the physical vestiges. Intellectual impairment ranges from minor learning disabilities to mental retardation. Many adults with FAS function at early grade-school level. They often lack social and communication skills and find it difficult to understand the consequences of actions, form friendships, take initiative, and interpret social cues.

Greek philosopher Aristotle noticed problems in children of alcoholic mothers more than 23 centuries ago. In the United States today, 1 to 3 of every 1,000 infants has the syndrome, meaning 2,000 to 6,000 affected children are born each year. Many more children have milder “alcohol-related effects.” A fetus of a woman with active alcoholism has a 30 to 45 percent chance of harm from prenatal alcohol exposure.

Nutrients

Certain nutrients ingested in large amounts, particularly vitamins, act as drugs. The acne medicine isotretinoin (Accutane) is a vitamin A derivative that causes spontaneous abortion and defects of the heart, nervous system, and face in exposed embryos. Physicians first noted the tragic effects of this drug 9 months after dermatologists began prescribing it to young women in the early 1980s. Another vitamin A-based drug, used to treat psoriasis, as well as excesses of vitamin A itself, also cause birth defects. Some forms of vitamin A are stored in body fat for up to 3 years.

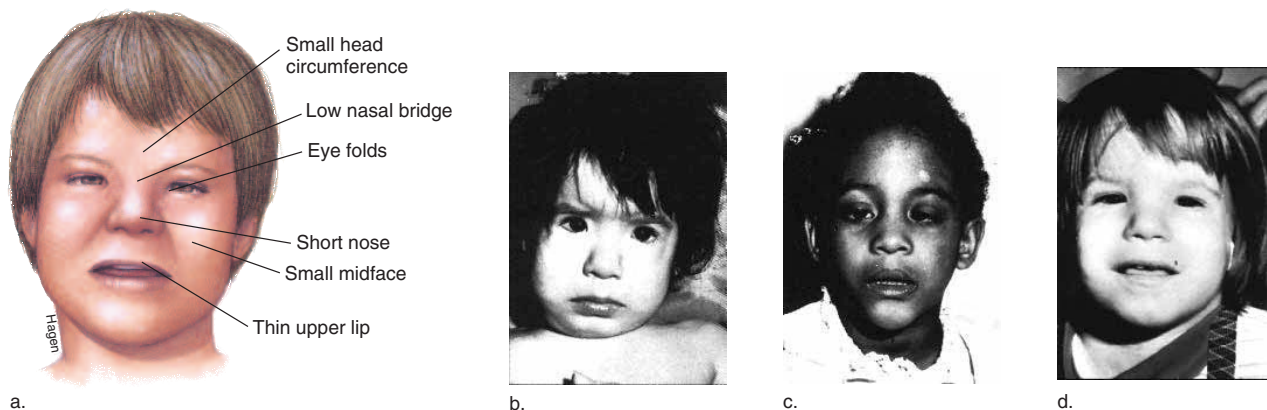


Figure 3.20 Fetal alcohol syndrome. Some children whose mothers drank alcohol during pregnancy have characteristic flat faces (a) that are strikingly similar in children of different races (b, c, and d).

Excess vitamin C can harm a fetus that becomes accustomed to the large amounts the woman takes. After birth, when the vitamin supply suddenly plummets, the baby may develop symptoms of vitamin C deficiency (scurvy), bruising and becoming infected easily.

Malnutrition threatens a fetus. Obstetrical records of pregnant women before, during, and after World War II link inadequate nutrition in early pregnancy to an increase in the incidence of spontaneous abortion. The aborted fetuses had very little brain tissue. People conceived during periods of famine who survived show ill effects, too. A study of people born just after the “Dutch Hunger Winter” of 1944–1945, for example, found that they suffered many more health problems throughout adulthood than their siblings who had adequate nutrition in the first 10 weeks of prenatal development. The individuals who were malnourished before birth also had different patterns of expression of key genes than their siblings. Poor nutrition later in pregnancy affects the development of the placenta and can cause low birth weight, short stature, tooth decay, delayed sexual development, and learning disabilities.

Occupational Hazards

Teratogens are present in some workplaces. Researchers note increased rates of spontaneous abortion and children born with birth defects among women who work with textile dyes, lead, certain photographic chemicals, semiconductor materials, mercury, and cadmium. Men whose jobs expose them to sustained heat, such as smelter workers, glass manufacturers, and bakers, may produce sperm that can fertilize an oocyte and then cause spontaneous abortion or a birth defect. A virus or a toxic chemical carried in semen may also cause a birth defect.

Viral Infection

Viruses are small enough to cross the placenta and reach a fetus. Some viruses that cause mild symptoms in an adult, such as the chickenpox virus, may devastate a fetus. Men can transmit viral infections to an embryo or fetus during sexual intercourse.

HIV can reach a fetus through the placenta or infect a newborn via blood contact during birth. Fifteen to 30 percent of infants born to untreated HIV-positive women are HIV positive. The risk of transmission is significantly reduced if a pregnant woman takes anti-HIV drugs. All fetuses of HIV-infected women are at higher risk for low birth weight, prematurity, and stillbirth if the woman’s health is failing.

German measles (rubella) is a well-known viral teratogen. In the United States, in the early 1960s, an epidemic of the usually mild illness caused 20,000 birth defects and 30,000 stillbirths. Children who were exposed during the first trimester of pregnancy could develop cataracts, deafness, and heart defects. Fetuses exposed during the second or third trimester of pregnancy may have as a result developed learning disabilities, speech and hearing problems, and type 1 diabetes mellitus.

The incidence of these problems, called congenital rubella syndrome, has dropped markedly since vaccination eliminated the disease in the United States. However, the syndrome resurfaces in unvaccinated populations. In 1991 among a cluster of unvaccinated

Amish women in rural Pennsylvania, 14 of every 1,000 newborns had congenital rubella syndrome, compared to an incidence then of 0.006 per 1,000 in the general U.S. population.

Herpes simplex virus can harm a fetus or newborn whose immune system is immature. Forty percent of babies exposed to active vaginal herpes lesions become infected, and half of them die. Of the survivors, 25 percent sustain severe nervous system damage, and another 25 percent have skin sores. A surgical (caesarean) delivery can protect the child.

Pregnant women are routinely checked for hepatitis B infection, which in adults causes liver inflammation, great fatigue, and other symptoms. Each year in the United States, 22,000 infants are infected with this virus during birth. These babies are healthy, but at high risk for developing serious liver problems as adults. When infected women are identified, a vaccine can be given to their newborns to help prevent complications.

Key Concepts

1. The critical period is the time during prenatal development when a structure is sensitive to damage from a faulty gene or environmental insult.
2. Most birth defects develop during the embryonic period and are more severe than problems that arise during fetal development.
3. Teratogens are agents that cause birth defects.

3.6 Maturation and Aging

“Aging” means moving through the life cycle. In adulthood, as we age, the limited life spans of cells are reflected in the waxing and waning of biological structures and functions. Although some aspects of our anatomy and physiology peak very early—such as the number of brain cells or hearing acuity, which do so in childhood—age 30 seems to be a turning point for decline. Some researchers estimate that, after this age, the human body becomes functionally less efficient by about 0.8 percent each year.

Many diseases that begin in adulthood, or are associated with aging, have genetic components. Often these disorders are multifactorial, because it takes many years for environmental exposures to alter gene expression in ways that noticeably affect health. Following is a closer look at how genes may impact health throughout life.

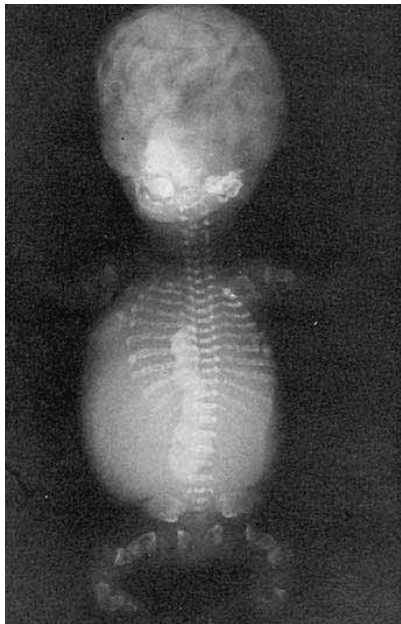
Adult-Onset Inherited Disorders

Human prenatal development is a highly regulated program of genetic switches that are turned on in specific body parts at specific times. Environmental factors can affect how certain genes are expressed before birth in ways that create risks that appear much later. For example, adaptations that enable a fetus to grow despite near-starvation become risk factors for certain common conditions of adulthood, when conserving calories is no longer

needed. Such disorders include coronary artery disease, obesity, stroke, hypertension, and type 2 diabetes mellitus. A fetus that does not receive adequate nutrition has intrauterine growth retardation (IUGR), and though born on time, is very small. Premature infants, in contrast, are small but are born early, and are not predisposed to conditions resulting from IUGR.

More than 100 studies clearly correlate low birth weight due to IUGR with increased incidence of cardiovascular disease later in life. Much of the data come from war records because enough time has elapsed to study the effects of prenatal malnutrition as people age. For example, a study of nearly 15,000 people born in Sweden from 1915 to 1929 correlates IUGR to heightened cardiovascular disease risk after age 65. Similarly, an analysis of individuals who were fetuses during a 7-month famine in the Netherlands in 1943 indicates a high rate of diabetes among them today. Experiments on intentionally starved sheep and rat fetuses support these historical findings.

How can poor nutrition before birth cause disease decades later? Perhaps to survive, the starving fetus redirects its circulation to protect vital organs such as the brain. At the same time, muscle mass and hormone production change to conserve energy. Growth-retarded babies have too little muscle tissue, and since muscle is the primary site of insulin action, glucose metabolism is altered. Thinness at birth, and the accelerated weight gain in childhood that often compensates, sets the stage for coronary heart disease and type 2 diabetes much later.



a.



b.

Figure 3.21 Genes act at various stages of development and life. (a) Osteogenesis imperfecta breaks bones, even before birth. This fetus has broken limb bones, a beaded appearance of the ribs due to fractures, and a poorly mineralized skull. (b) At the funeral of former president Richard M. Nixon in April 1994, all was not right with former president Ronald Reagan. He was forgetful and responded inappropriately to questions. Six months later he penned a moving letter confirming that he had Alzheimer disease. By 1997, Reagan no longer knew the names of his closest relatives. By 1999, he didn't remember anyone, and by 2001 he no longer recalled being president. He died in June 2004.

In contrast to the delayed effects of fetal malnutrition, symptoms of single-gene disorders can begin at any time. In general, inherited conditions that affect children are recessive. Even a fetus can have symptoms of inherited disease, as **figure 3.21a** illustrates for osteogenesis imperfecta (“brittle bone disease”, MIM 166210).

Most dominantly inherited conditions start to affect health in early to middle adulthood. This is the case for polycystic kidney disease (MIM 173900). Cysts that may have been present but undetected in the kidneys during one's twenties begin causing bloody urine, high blood pressure, and abdominal pain in the thirties. Similarly, the joint destruction of osteoarthritis may begin in one's thirties, but not cause pain for 20 years. The uncontrollable movements, unsteady gait, and diminishing mental faculties of Huntington disease typically become obvious near age 40, but may have begun years earlier.

Five to 10 percent of Alzheimer disease cases are inherited and produce initial symptoms in the forties and fifties. German neurologist Alois Alzheimer first identified the condition in 1907 as affecting people in mid-adulthood. Noninherited Alzheimer disease typically begins later in life (**figure 3.21b**).

Whatever the age of onset, Alzheimer disease starts gradually. Mental function declines steadily for 3 to 10 years after the first symptoms of depression and short-term memory loss appear. The person gradually loses cognitive skills, even becoming unable to walk because he or she forgets how to put one foot in front of the other. Confused and forgetful, Alzheimer patients often wander away from family and friends. Finally, the patient cannot recognize loved ones and can no longer perform basic functions such as speaking or eating.

The brains of Alzheimer disease patients contain deposits of a protein called beta amyloid in learning and memory centers. Alzheimer brains also contain structures called neurofibrillary tangles, which consist of a protein called tau. Tau binds to and disrupts microtubules in nerve cell branches, destroying the shape of the cell, which is essential to its ability to communicate. It isn't clear whether the “plaques and tangles” of the Alzheimer brain are causes or effects. Reading 5.1 discusses Alzheimer disease further.

Disorders That Resemble Accelerated Aging

Genes control aging both passively (as structures break down) and actively (by initiating new activities). A group of “rapid aging” inherited disorders may hold clues to how genes control aging in all of us.

The most severe rapid aging disorders are the segmental progeroid syndromes. (They were once called progerias, but the newer terminology reflects

the fact that they do not hasten all aspects of aging.) Most of these disorders are caused by impairment of cells' ability to repair DNA, which is discussed in chapter 12. Mutations that would ordinarily be corrected persist. Over time, the accumulation of mutations destabilizes the entire genome, and even more mutations occur in somatic cells. The various changes that we associate with aging ensue.

Table 3.3 lists the more common segmental progeroid syndromes. They vary in severity. People with Rothmund-Thomson syndrome, for example, may have a normal life span, but develop gray hair or baldness, cataracts, cancers, and osteoporosis at young ages. Werner syndrome becomes apparent before age 20, causing death before age 50 from diseases associated with aging. Young adults with Werner syndrome develop atherosclerosis, type 2 diabetes mellitus, hair graying and loss, osteoporosis, cataracts, and wrinkled skin. They are short because they skip the growth spurt of adolescence.

The child in **figure 3.22**, Megan, has Hutchinson-Gilford progeria syndrome, but is unusually beautiful. An affected child typically appears normal at birth but slows in growth by the first birthday. Within just a few years, the child becomes wrinkled and bald, with easily bruised skin and the facial features characteristic of advanced age. The body ages on the inside as well, as arteries clog with fatty deposits. The child usually dies of a heart attack or a stroke by age 13, although some patients live into their twenties.

The child's cells show aging-related changes too. Normal cells growing in culture divide about 50 times before dying. Cells from Hutchinson-Gilford progeria syndrome patients die in culture after only 10 to 30 divisions.

Only 50 people who currently have Hutchinson-Gilford progeria syndrome are known worldwide, and only 100 have ever been reported in the medical literature. Identifying the gene mutation in 2003 led to a key understanding about the molecular cause and identification of drugs that might help treat this fatal disease. From 2005 to 2006, a team of researchers studied 15 patients, to better describe the phenotype. Unlocking the molecular cause of the disease has suggested several uncommon drugs (farnesyltransferase inhibitors) and several commonly used drugs (statins and bisphosphonates) that are now in clinical trials to help stall the symptoms.



Figure 3.22 Accelerated aging. Megan has Hutchinson-Gilford progeria syndrome, which is extremely rare.

The disease is caused by a single DNA base change in the gene that encodes a protein called lamin A. That one base is a site that determines how parts of the protein are cut and joined, and when it is altered, the protein lacks 50 amino acids. The shortened protein is called progerin. It remains stuck to the endoplasmic reticulum, instead of being transported into the nucleus through the nuclear pores, as happens to the normal lamin A protein. Instead, progerin diffuses into the tubules of the ER and travels within them to the nuclear membrane, which is possible because these two organelles are continuous. This route of entry stresses the nuclear membrane, causing it to bubble or “bleb” inward, altering the way that the nuclear lamina (the layer on the inside face) binds the chromatin (DNA complexed with protein) within. Somehow, disturbing the chromatin hampers DNA repair, allowing mutations associated with the signs of aging to occur. The drugs being tested block the molecule that holds progerin to the ER.

Studies on stem cells from bone marrow of patients add further evidence: Progerin shifts the activities of certain genes in ways that promote bone formation and suppress fat deposition. This skewed development perhaps explains the failure to thrive and skeletal appearance of affected individuals. The molecular view of progeria suggests that DNA repair is what enables us to live many years. These children lack that protection, and the mutations that age us all accumulate much faster.

Table 3.3 Rapid Aging Syndromes

Disorder	Incidence	Average Life Span	MIM Number
Ataxia telangiectasia	1/60,000	20	208900
Cockayne syndrome	1/100,000	20	216400
Hutchinson-Gilford syndrome	<1/1,000,000	13	176670
Rothmund-Thomson syndrome	<1/100,000	normal	268400
Trichothiodystrophy	<1/100,000	10	601675
Werner syndrome	<1/100,000	50	277700

Is Longevity Inherited?

Aging reflects genetic activity plus a lifetime of environmental influences. Families with many very aged members have a fortuitous collection of genes plus shared environmental influences such as good nutrition, excellent health care, devoted relatives, and other advantages. A genome-level approach to identifying causes of longevity identified a region of chromosome 4 that houses gene variants associated with long life. Genome comparisons among people who've passed their 100th birthdays and those who have died of the common illnesses of older age will reveal other genes that influence longevity (**Reading 3.1**).

It is difficult to tease apart inborn from environmental influences on life span. One approach compares adopted individuals to both their biological and adoptive parents. In a study from Denmark, adopted individuals with one biological parent who died of natural causes before age 50 were more than twice as likely to die before age 50 as were adoptees whose biological parents lived beyond this age, suggesting an inherited component to longevity. Interestingly, adopted individuals whose natural parents died early due to infection were more than five times as likely to also die early of infection, perhaps because

of inherited immune system deficiencies. The adoptive parents' ages at death had no influence on that of their adopted children. Chapter 7 explores the "nature versus nurture" phenomenon more closely.

Key Concepts

1. Starvation before birth can set the stage for later disease by affecting gene expression in certain ways.
2. Most single-gene disorders are recessive and strike early in life. Single-gene disorders with an adult onset are more likely to be dominant.
3. The segmental progeroid syndromes are single-gene disorders that speed the signs of aging.
4. Families with many aged members can thank their genes as well as the environment. Chromosome 4 houses longevity genes, and genome-wide screens are identifying others.
5. Adoption studies compare the effects of genes versus environmental influences on longevity.

Reading 3.1

Genes and Longevity

The human genome is like a vast library that holds the clues to good health. One way to identify those clues is to probe the genomes of individuals who have lived past 100 years. These fortunate people are called centenarians (**figure 1**). Usually they enjoy excellent health and are socially active, then succumb rapidly to diseases that typically claim people decades earlier.

Centenarians fall into three broad groups—about 20 percent of them never get the diseases that kill most people; 40 percent get these diseases, but at a much older age than average; and the other 40 percent live with and survive the more common disorders of aging. Researchers hope that learning which gene variants offer this protection will lead to better understanding of the common disorders of later adulthood—heart disease, stroke, cancers, type 2 diabetes mellitus, and dementias.

While the environment seems to play an important role in the deaths of people ages 60 to 85, past that age, genes predominate. That is, someone who dies at age 68 of lung cancer can probably blame a lifetime of cigarette smoking. But a smoker who dies at age 101 of the same disease probably had gene variants that protected against lung cancer. Centenarians have higher levels of large lipoproteins that carry cholesterol (HDL) than other people, which researchers estimate adds 20 years of life.

Children and siblings of centenarians tend to be long-lived as well, supporting the idea that longevity is inherited. Brothers of centenarians are 17 times as likely to live past age 100 as the average



Figure 1 This woman has enjoyed living for more than a century. Researchers are discovering clues to good health by probing the genomes of centenarians.

(Continued)

(Continued)

man, and sisters are 8.5 times as likely. The fact that some people more than 100 years of age have less-than-healthy habits suggests that genes are protecting them. One researcher suggests that the saying, “The older you get, the sicker you get” be replaced with “The older you get, the healthier you’ve been.”

Centenarians have inherited two types of gene variants—those that directly protect them, and wild type alleles of genes that, when mutant, cause disease. Research focuses on individual genes as well as genome-wide scans to identify gene variants that make it more likely that a person will live past age 100. **Table 1** lists some “candidate” gene types that may control longevity. To find other gene variants that promote long life, researchers are conducting genome-wide association studies on centenarians and comparing the patterns to those of people with particular conditions associated with aging.

Several studies are identifying gene variants that contribute to living long and well. The New England Centenarian Study, headed at Boston University, began in 1988 to amass information on families of the oldest citizens in the United States. The researchers are compiling a “healthy standard genome.” Investigators from the Coriell Institute in New Jersey are probing the genomes of people over age 90 who live in nursing homes. So far, what these people have in common is never having had heart disease and never having smoked. Several had cancer, indicating that cancers are often survivable.

Researchers at the University of Pittsburgh have identified places in the genome that harbor “successful aging genes” that have variants that preserve cognition. Other studies are looking at known genes implicated in the diseases that kill most of us, while others seek patterns of variation or the alleles that long-lived siblings share. Considered together, perhaps these studies will provide information that will help the majority of us who have not been fortunate enough to have inherited longevity gene variants.

Table 1

Single genes important in aging affect

- control of insulin secretion and glucose metabolism,
- immune system functioning,
- control of the cell cycle,
- lipid (cholesterol) metabolism,
- response to stress, and
- production of antioxidant enzymes.

Summary

3.1 The Reproductive System

1. The male and female reproductive systems include paired **gonads** and networks of tubes in which **sperm** and **oocytes** are manufactured.
2. Male **gametes** originate in seminiferous tubules within the paired testes. They then pass through the epididymis and ductus deferentia, where they mature before exiting the body through the urethra during sexual intercourse. The prostate gland, the seminal vesicles, and the bulbourethral glands add secretions.
3. Female gametes originate in the ovaries. Each month after puberty, one ovary releases an oocyte into a uterine tube. The oocyte then moves to the uterus for implantation (if fertilized) or expulsion.

3.2 Meiosis

4. **Meiosis** reduces the chromosome number in gametes from **diploid** to **haploid**, maintaining the chromosome number between generations. Meiosis ensures genetic variability by **independently assorting** combinations of genes into gametes as chromosomes randomly align and **cross over**.
5. Meiosis I, a **reduction division**, halves the number of chromosomes. Meiosis II, an **equational division**, produces four cells from the two that result from meiosis I, without another DNA replication.
6. Crossing over occurs during prophase I. It mixes up paternally and maternally derived genes on **homologous pairs** of chromosomes.
7. Chromosomes segregate and independently assort in metaphase I, which determines the distribution of genes from each parent.

3.3 Gamete Maturation

8. **Spermatogenesis** begins with spermatogonia, which accumulate cytoplasm and replicate their DNA, becoming primary spermatocytes. After meiosis I, the cells become haploid secondary spermatocytes. In meiosis II, the secondary spermatocytes divide, each yielding two spermatids, which then differentiate into spermatozoa.
9. In **oogenesis**, some oogonia grow and replicate their DNA, becoming primary oocytes. In meiosis I, the primary oocyte divides to yield one large secondary oocyte and a small **polar body**. In meiosis II, the secondary oocyte divides to yield the large ovum and another polar body. Female meiosis is completed at fertilization.

3.4 Prenatal Development

10. In the female, sperm are capacitated and drawn toward a secondary oocyte. One sperm burrows through the oocyte’s protective layers with acrosomal enzymes. Fertilization occurs when the sperm and oocyte fuse and their DNA combines in one nucleus, forming the **zygote**. Electrochemical changes in the egg surface block additional sperm from entering. **Cleavage** begins and a 16-celled **morula** forms. Between days 3 and 6, the morula arrives at the uterus and hollows, forming a **blastocyst** made up of **blastomeres**. The trophoblast and **inner cell mass** form. Around day 6 or 7, the blastocyst implants, and trophoblast cells secrete hCG, which prevents menstruation.
11. During the second week, the amniotic cavity forms as the inner cell mass flattens. **Ectoderm** and **endoderm** form, and then **mesoderm** appears, establishing the **primary germ layers**. Cells in each germ layer begin to develop into specific

organs. During the third week, the placenta, yolk sac, allantois, and umbilical cord begin to form as the amniotic cavity swells with fluid. **Monozygotic** twins result when one fertilized ovum splits. **Dizygotic** twins result from two fertilized ova. Organs form throughout the embryonic period. The primitive streak, notochord and **neural tube**, arm and leg buds, heart, facial features, and skeleton develop.

12. At the eighth week, the **embryo** becomes a **fetus**, with all structures present but not fully grown. Organ rudiments grow and specialize. The developing organism moves and reacts, and its body proportions come to resemble those of a baby. In the last trimester, the brain develops rapidly, and fat is deposited beneath the skin. The digestive and respiratory systems mature last.

3.5 Birth Defects

13. Birth defects can result from a mutation or an environmental intervention.

14. A substance that causes birth defects is a **teratogen**. Environmentally caused birth defects are not transmitted to future generations.
15. The time when a structure is sensitive to damage from an abnormal gene or environmental intervention is its **critical period**.

3.6 Maturation and Aging

16. Genes cause or predispose us to illness throughout life. Single-gene disorders that strike early tend to be recessive; most adult-onset single-gene conditions are dominant.
17. Malnutrition before birth can alter gene expression in ways that cause illness later in life.
18. The segmental progeroid syndromes are single-gene disorders that speed aging-associated changes.
19. Long life is due to genetics and environmental influences.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

- How many sets of human chromosomes are in each of the following cell types?
 - an oogonium
 - a primary spermatocyte
 - a spermatid
 - a cell from either sex during anaphase of meiosis I
 - a cell from either sex during anaphase of meiosis II
 - a secondary oocyte
 - a polar body derived from a primary oocyte
- List the structures and functions of the male and female reproductive systems.
- A dog has 39 pairs of chromosomes. Considering only the random alignment of chromosomes, how many genetically different puppies are possible when two dogs mate? Is this number an underestimate or overestimate of the actual total? Why?
- How does meiosis differ from mitosis?
- What do oogenesis and spermatogenesis have in common, and how do they differ?
- How does gamete maturation differ in the male and female?
- Why is it necessary for spermatogenesis and oogenesis to generate stem cells?
- Describe the events of fertilization.
- Write the time sequence in which the following structures begin to develop: notochord, gastrula, inner cell mass, fetus, zygote, morula.
- Why does exposure to teratogens produce more severe health effects in an embryo than in a fetus?
- The same birth defect syndrome can be caused by a mutant gene or exposure to a teratogen. How do the consequences of each cause differ for future generations?
- List four teratogens, and explain how they disrupt prenatal development.
- Why is an "anti-aging" pill, diet, or device impossible?
- Cite two pieces of evidence that genes control aging.

Applied Questions

- Up to what stage, if any, do you think it is ethical to experiment on a prenatal human? Cite reasons for your answer.
- Under a microscope, a first and second polar body look alike. What structure would distinguish them?
- Armadillos always give birth to identical quadruplets. Are the offspring clones?
- Some Vietnam War veterans who were exposed to the herbicide Agent Orange claim that their children—born years after the exposure—have birth defects caused by dioxin, a contaminant in the herbicide. What types of cells would the chemical have to have affected in these men to cause birth defects years later?

5. In about 1 in 200 pregnancies, a sperm fertilizes a polar body instead of an oocyte. A mass of tissue that is not an embryo develops. Why can't a polar body support the development of an entire embryo?
6. Should a woman be held legally responsible if she drinks alcohol, smokes, or abuses drugs during pregnancy and it harms her child? Should liability apply to all substances that can harm a fetus, or only to those that are illegal?
7. Would you want to have your genome scanned to estimate how long you are likely to live? Why or why not?
8. What types of evidence have led researchers to hypothesize that a poor prenatal environment can raise the risk for certain adult illnesses? How are genes part of this picture?

Web Activities

9. Look over the "Living to 100 Life Expectancy Calculator" at www.livingto100.com and list 10 ways that you can change your behavior to possibly live longer. What does this quiz suggest about the relative role of genes and the environment in determining longevity?
10. Go to the Motherisk website at <http://www.motherisk.org/>. Click on "Women, Partners, Family and Friends." Then click on "drugs in pregnancy" in the left-hand list. Identify three drugs that are safe to take during pregnancy and three that are not safe, and list the associated medical problems.

Case Studies and Research Results

11. Human embryonic stem cells can be derived and cultured from an 8-celled cleavage embryo and from a cell of an inner cell mass. Explain the difference between these stages of human prenatal development.
12. Victor, a 34-year-old artist, was killed in a car accident. He and his wife Emma hadn't started a family yet, but planned to soon. The morning after the accident, Emma asked if some of her husband's sperm could be collected and frozen, for her to use to have a child. Do you think that this "post-mortem sperm retrieval" should be done? See the chapter opener for chapter 21 for further information.
13. Miguel and Maria are carriers of cystic fibrosis (CF), and the condition is severe in their families. They have a procedure called preimplantation genetic diagnosis (see figure 21.5) to ensure that they conceive a child who does not inherit CF. Maria's oocyte is fertilized with Miguel's sperm in a laboratory dish, and it develops to the 8-cell stage. One cell is removed and tested for the mutant allele that Miguel and Maria carry. Only the wild type allele is detected.

Anna and Peter are also carriers of a genetic disorder that can affect either sex. They cannot get into a preimplantation genetic diagnosis clinical trial, which would be free; their insurance will not cover the procedure; and they cannot afford it. So, they choose chorionic villus sampling (CVS), in which a cell from the developing placenta is tested for the mutant allele, at the tenth week of gestation. Their fetus is found to be a carrier, like them.

A third couple, Vivian and Max, are not willing to take the higher risk of miscarriage associated with (CVS), so they wait until the sixteenth week, and Vivian has amniocentesis. Fetal cells are sampled from the amniotic fluid and the mutation that causes the clotting disorder hemophilia in Vivian's family checked. Vivian may be a carrier for hemophilia A. If she is, a son would face a 50 percent chance of inheriting the disorder. The amniocentesis indicates a daughter.

 - a. Why can a genetic test on a cell from an embryo, the placenta, or a fetus predict future health?
 - b. At the time of preimplantation genetic diagnosis, is the embryo a cleavage embryo, an inner cell mass, or a gastrula?
 - c. What structures are present in Vivian and Max's fetus that have not yet developed in Anna and Peter's at the time of their prenatal tests?
14. Surgical separation of conjoined twins is more likely to succeed if fewer body parts are shared or attached. This was the case for Maria de Jesus and Maria Theresa, born in Guatemala in 2001 and separated before their first birthday. They were joined at the head, but facing opposite directions, so they could not move much. The surgery took 23 hours! Today they are well. The outcome wasn't good for Mandan and Label Bikini, 29-year-old Iranian conjoined twins who could no longer stand being joined along their heads, with their brains fused. They died shortly after 50 hours of surgery in 2003.

If you had conjoined twins, what would you do? Would you attempt surgical separation?

PART 2 Transmission Genetics



Bea Rienhoff has a genetic disease, but it is so rare that so far, it hasn't been diagnosed. Her father, a physician and geneticist, sequenced Bea's DNA himself, searching for an answer. His website, mydaughtersdna.org, has helped others understand their symptoms.

CHAPTER

4

Single-Gene
Inheritance

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His Daughter's DNA

When Hugh Rienhoff first saw his daughter Bea, he knew something wasn't quite right. An internist and clinical geneticist, Dr. Rienhoff took in the long feet, clenched fingers, "floppy" muscles, and facial birthmark and suspected a single-gene disorder. Such conditions, usually extremely rare, affect many organ systems and produce peculiar telltale signs, such as the slant of the eyes or the spacing between the nose and the upper lip. This constellation of findings is called a syndrome, which is Greek for "running together." Unless a physician considers the entire patient, a diagnosis can easily be missed. But some disorders are so rare that they do not even have a name, let alone a description. That would turn out to be the case with Hugh Rienhoff's daughter.

As Bea got older but failed to gain much weight, appearing thin with birdlike legs, her father recognized some signs of Marfan syndrome (MIM 154700) and some of Loey's-Dietz syndrome (MIM 609192, 610380). But since pediatricians could not definitively diagnose her, Dr. Rienhoff took matters into his own hands. He sequenced specific genes in Bea's DNA himself, genes that he thought might account for the symptoms. "It was eerie examining her DNA, as though I were peering through a powerful microscope looking deep into my daughter while she patiently lay on the microscope stage, looking up, hoping for answers."

Dr. Rienhoff started a website, mydaughtersdna.org, where others looking for diagnoses could post test results and descriptions, to see if symptoms fit a known pattern. Although a gene responsible for Bea's condition has yet to be found, at least one person now knows her own diagnosis, thanks to the website. Wrote Dr. Rienhoff, "Though historically genetics has been focused

on the rare and unusual, and dominated by a handful of experts, the new wave of understanding of the human genome cannot happen without our large-scale participation. . . . each of us has something very important to contribute to human genomics research: our DNA and our phenotype. It is perhaps the most profound experiment humankind will undertake to understand itself as a species." The National Institutes of Health has begun a program to discover the causes of "one-of-a-kind" conditions such as the one that affects young Bea Rienhoff.

4.1 A Tale of Two Families

Henry T. is a healthy 21-year-old college student trying to decide whether or not to take a genetic test that will reveal if he has inherited the mutation that causes Huntington disease (HD). He has been meeting with a genetic counselor, but he doesn't need a medical professional to educate him about HD—he's seen it all his life. **Figure 4.1** is a timeline/narrative of the effects of HD on Henry's family.

Henry doesn't remember much about his grandfather Emmett, who died last year. Emmett had been in a nursing

home for as long as Henry can recall, just down the hall from his younger sister, great aunt Louise. When Henry turned 12, his parents told him about the family illness and took him to visit his grandfather and great aunt. Their descriptions hadn't prepared him for the sight of his grandfather restrained in bed by tightly tucked blankets that could barely quell his constant writhing. His face grimaced and ticked, and he had great difficulty speaking. By the time Henry entered college, his grandfather had become curiously still, as if he had run out of energy after years of near-constant moving.

Unfortunately, HD didn't leave Henry's family when his grandfather died—his mother Ann was beginning to show subtle symptoms. She moved her left cheek in a funny way, repeatedly. Something was not quite right about her eye movements. She'd yell at whoever was around, blowing everyday things way out of proportion. On Thanksgiving morning Henry found his mother sitting in the middle of the kitchen floor, crying, surrounded by the mess of a half-cooked turkey. He was worried—about his mother, but also about himself and his younger sister. For he knew that HD does not skip generations, unless, by chance, no one inherits the mutation.

Sean P., like Henry, doesn't actually have symptoms of a genetic disease, but he has lived with one in his family all his life. Although he and his parents are healthy, his younger sister

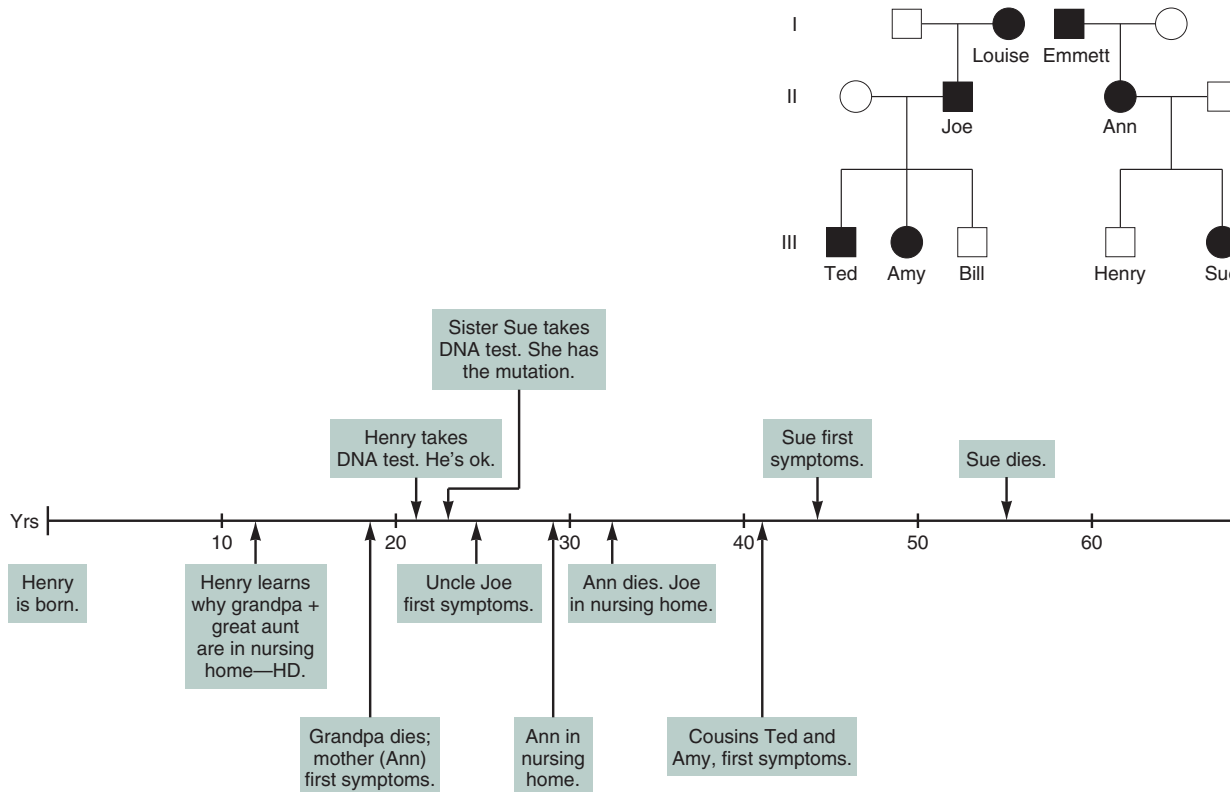


Figure 4.1 An inherited disease timeline. Huntington disease reverberates through a family, striking each generation until all affected individuals do not have children who have inherited the mutation. Henry lives with survivor guilt that he is unaffected. He and his cousin Bill help each other to care for their affected parents, and then their affected siblings. The inset shows a pedigree. These diagrams are discussed later in the chapter, but the inheritance pattern is clear in the filled-in symbols.

Ellen has cystic fibrosis (CF). Sean remembers looking forward to the arrival of the baby when he was in kindergarten, but never expected the extent of her health problems.

Ellen couldn't gain weight, and she seemed always to have respiratory infections and difficulty breathing. When bronchitis turned into pneumonia when she was only 8 months old, her pediatrician ordered a test for CF—Ellen indeed had a classic case, with two copies of the most common mutation. That was several years ago. Today, Sean's mother's carrier status would have been picked up during pregnancy, and then his father tested, and he, too, would have been found to be a carrier. Then newborn screening would have revealed that Ellen had inherited the mutation from both parents, and treatment begun—enzymes to help digestion and breathing, and antibiotics to prevent infection.

The greatest impact on the children was the twice-daily “airway clearance” respiratory therapy that the parents took turns performing. At first Sean thought that they were punishing his baby sister when they turned her in different positions and hit her back, but quickly realized that they were shaking free the thick mucus that was clogging her lungs (figure 4.2).

Ellen's childhood was rough, with frequent hospital stays and constant vigilance to keep her free of infection. No one else in the family had the disease. One summer, Sean went to a summer camp for CF families, and it was there that he realized that he could be a carrier, like his parents. So, a few years later, he took a CF test at the campus health center, and learned that he, too, is a carrier. Reading 4.2 at the end of the chapter has further information on CF.

Huntington disease and cystic fibrosis affect families in patterns termed **modes of inheritance**. Knowing these patterns makes it possible to predict risks that particular family members inherit the mutation. HD is **autosomal dominant**, which means that it affects both sexes and appears every generation. In contrast, CF is **autosomal recessive**, which means that it affects both sexes and can skip generations through carriers. Henry's sister became ill because only one mutant gene is necessary for this to happen in an autosomal dominant disease. Sean will *not* become ill because two mutations are required to inherit an autosomal recessive disease.



Figure 4.2 Treating cystic fibrosis. In cystic fibrosis, the thick, sticky mucus that clogs airways must be coughed up at least twice every day.

This chapter explains the roots of the modes of inheritance. The tale begins with pea plants in a long-ago garden in a small village in what is now the Czech Republic.

4.2 Following the Inheritance of One Gene—Segregation

Today we think of genetics in terms of traits and health, but the field actually grew out of agriculture, which focuses on breeding certain plants and animals for traits that we value. Gregor Mendel was the first thinker to probe the underlying rules of logic that make it possible to predict inheritance of specific traits. In a series of clever breeding experiments in pea plants, Mendel described units of inheritance that pass traits from generation to generation. He called these units “elementen.” He could not see them, but he inferred their existence from the appearances of his plants. Although Mendel knew nothing of DNA, chromosomes, or cells, his “laws” of inheritance have not only stood the test of time, but explain trait transmission in any diploid species.

Mendel the Man

The son of a farmer and grandson of a gardener, Mendel learned early how to tend fruit trees. At age 10 he left home to attend a special school for bright students, supporting himself by tutoring. He eventually became a priest at an atypical Augustinian monastery where the priests were teachers and did research in natural science. Here, Mendel learned how to artificially pollinate crop plants to control their breeding.

Mendel wanted to teach natural history, but had difficulty passing his exams due to test anxiety. At age 29, he was such an effective substitute teacher that he was sent to earn a college degree. At the University of Vienna, courses in the sciences and statistics fueled his interest in plant breeding and got him thinking about a question that had confounded other plant breeders—why did certain traits disappear in one generation, yet reappear in the next? To solve this puzzle, Mendel bred pea plant hybrids and applied statistics.

From 1857 to 1863, Mendel crossed and cataloged traits in 24,034 plants, through several generations. He deduced that consistent ratios of traits in the offspring indicated that the plants transmitted distinct units. He derived two hypotheses to explain how inherited traits are transmitted. Mendel described his work to the Brnő Medical Society in 1865 and published it in the organization's journal the next year. The remarkably clear paper discussed plant hybridization, the reappearance of traits in the third generation, and the joys of working with peas, plus data.

It took years for Mendel's findings to be recognized. Shortly after his treatise was published in English in 1901, three botanists (Hugo DeVries, Karl Franz Joseph Erich Correns, and Seysenegg Tschermak) independently rediscovered the laws of inheritance. Once they read Mendel's paper, they credited him. Mendel came to be regarded as the “father of genetics.”

In the twentieth century, researchers discovered the molecular basis of some of the traits that Mendel studied. “Short” and “tall” plants reflect expression of a gene that enables a plant to

produce the hormone gibberellin, which elongates the stem. One tiny change to the DNA, and a short plant results. Likewise, “round” and “wrinkled” peas arise from the *R* gene, whose encoded protein connects sugars into branching polysaccharide molecules. Seeds with a mutant *R* gene cannot attach the sugars. As a result, water exits the cells, and the peas wrinkle.

Mendel’s Experiments

Peas are ideal for probing heredity because they are easy to grow, develop quickly, and have many traits that take one of two easily distinguishable forms. **Figure 4.3** illustrates the seven traits that Mendel followed through several pea generations. When analyzing genetic crosses, the first generation is the parental generation, or P_1 ; the second generation is the first filial generation, or F_1 ; the next generation is the second filial generation, or F_2 , and so on.

Mendel’s first experiments dealt with single traits with two expressions, such as “short” and “tall.” He set up all combinations of possible artificial pollinations, manipulating fertilizations to cross tall with tall, short with short, and tall with short plants. This last combination, plants with one trait variant crossed to plants with the alternate, produces hybrids, which are offspring that inherit a different gene variant from each parent.

Mendel noted that short plants crossed to other short plants were “true-breeding,” always producing short plants. The crosses of tall plants to each other were more confusing. Some tall plants were true-breeding, but others crossed with each other yielded short plants in about one-quarter of the next generation. It appeared as if in some tall plants, tallness could mask shortness. One trait that masks another is said to be **dominant**; the masked trait is **recessive**.

Mendel conducted up to 70 hybrid crosses for each of the seven traits. Because one trait is followed and the parents are hybrids, this is called a **monohybrid cross**.

When Mendel allowed the non-true-breeding tall plants—monohybrids—to self-fertilize, the progeny were in the ratio of one-quarter short to three-quarters tall plants (**figure 4.4**). In further crosses, he found that two-thirds of the tall plants from the monohybrid F_1 cross were non-true-breeding, and the remaining third were true-breeding.

In these experiments, Mendel confirmed that hybrids hide one expression of a trait—short, in this case—which reappears when hybrids are self-crossed. He tried to explain how this happened: Gametes distribute “elementen” because these cells physically link generations. Paired sets of elementen separate as gametes form. When gametes join at fertilization, the elementen combine anew. Mendel reasoned that each elementen was packaged in a separate gamete. If opposite-sex gametes combine at random, he could mathematically explain the different ratios of traits produced from his pea plant crosses. Mendel’s idea that elementen separate in the gametes would later be called the **law of segregation**.

When Mendel’s ratios were demonstrated in several species in the early 1900s, just when chromosomes were being discovered, it became apparent that elementen and chromosomes had much in common. Both paired elementen and pairs of chromosomes separate at each generation and are transmitted—one from each parent—to offspring. Both are inherited in random combinations. Chromosomes provided a physical mechanism for Mendel’s hypotheses. In 1909, English embryologist William Bateson renamed Mendel’s elementen *genes* (Greek for “give birth to”). In the 1940s, scientists began investigating the gene’s chemical basis, discussed in chapter 9.















	Seed form	Seed color	Pod form	Pod color	Flower position	Seed coat color	Stem length
Dominant	 Round (<i>R</i>)	 Yellow (<i>Y</i>)	 Inflated (<i>V</i>)	 Green (<i>G</i>)	 Axial (<i>F</i>) (along stem)	 Gray or gray-brown (<i>A</i>)	 Tall (<i>T</i>)
Recessive	 Wrinkled (<i>r</i>)	 Green (<i>y</i>)	 Restricted (<i>v</i>)	 Yellow (<i>g</i>)	 Terminal (<i>f</i>) (on top)	 White (<i>a</i>)	 Short (<i>t</i>)

Figure 4.3 Traits Mendel studied. Gregor Mendel studied the transmission of seven traits in the pea plant. Each trait has two easily distinguished expressions, or phenotypes.

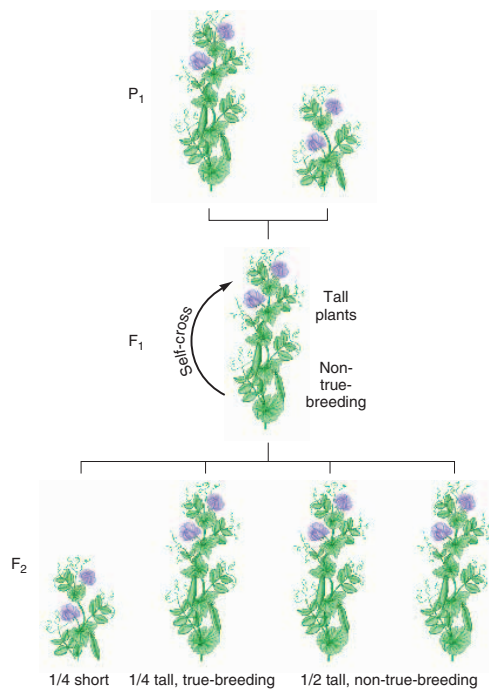


Figure 4.4 A monohybrid cross. When Mendel crossed true-breeding tall plants with short plants, the next generation plants were all tall. When he self-crossed the F₁ plants, one-quarter of the plants in the next generation, the F₂, were short, and three-quarters were tall. Of the tall plants in the F₂, one-third were true-breeding, and the other two-thirds were not true-breeding. He could tell this by conducting further crosses of the tall plants to short plants, to see which bred true.

Terms and Tools to Follow Segregating Genes

The law of segregation reflects the actions of chromosomes and genes during meiosis. Because a gene is a long sequence of DNA, it can vary in many ways. An individual with two identical alleles for a gene is **homozygous** for that gene. An individual with two different alleles is **heterozygous**—what Mendel called “non-true-breeding” or “hybrid.”

When considering a gene with two alleles, the dominant one is shown as a capital letter and the recessive with the corresponding small letter. If both alleles are recessive, the individual is homozygous recessive, shown with two small letters. An individual with two dominant alleles is homozygous dominant, and has two capital letters. One dominant and one recessive allele, such as *Tt* for non-true-breeding tall pea plants, indicates heterozygotes.

An organism’s appearance does not always reveal its alleles. Both a *TT* and a *Tt* pea plant are tall, but *TT* is a homozygote and *Tt* a heterozygote. The **genotype** describes the organism’s alleles, and the **phenotype** describes the outward expression of an allele combination. A **wild type** phenotype is the most common expression of a particular allele combination in a population. The wild type allele may be recessive or dominant. A **mutant** phenotype is a variant of a gene’s expression that arises when the gene undergoes a change, or **mutation**.

Mendel was observing the events of meiosis. When a gamete is produced, the two copies of a gene separate with the homologs that carry them. In a plant of genotype *Tt*, for example, gametes carrying either *T* or *t* form in equal numbers during anaphase I. Gametes combine at random. A *t*-bearing oocyte is neither more nor less attractive to a sperm than is a *T*-bearing oocyte. These two factors—equal allele distribution into gametes and random combinations of gametes—underlie Mendel’s law of segregation (**figure 4.5**).

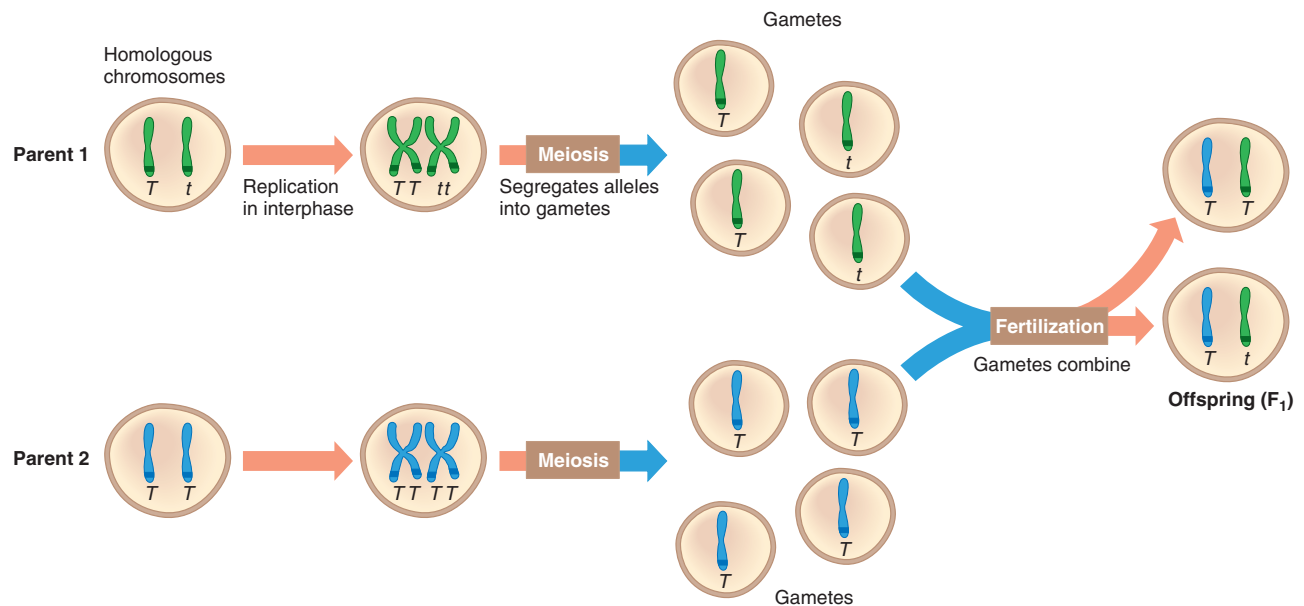


Figure 4.5 Mendel’s first law—gene segregation. During meiosis, homologous pairs of chromosomes and their genes separate and are packaged into separate gametes. At fertilization, gametes combine at random. Green and blue denote different parental origins of the chromosomes. This cross yields offspring of genotypes *TT* and *Tt*.

When Mendel crossed short plants (tt) with true-breeding tall plants (TT), the seeds grew into F_1 plants that were all tall (genotype Tt). Next, he self-crossed the F_1 plants. The progeny were TT , tt , and Tt . A TT individual resulted when a T sperm fertilized a T oocyte; a tt plant resulted when a t oocyte met a t sperm; and a Tt individual resulted when either a t sperm fertilized a T oocyte, or a T sperm fertilized a t oocyte.

Because two of the four possible gamete combinations produce a heterozygote, and each of the others produces a homozygote, the genotypic ratio expected of a monohybrid cross is $1TT: 2Tt: 1tt$. The corresponding phenotypic ratio is three tall plants to one short plant, a 3:1 ratio. Mendel saw these results for all seven traits that he studied (table 4.1). A diagram called a **Punnett square** shows these ratios (figure 4.6). A Punnett

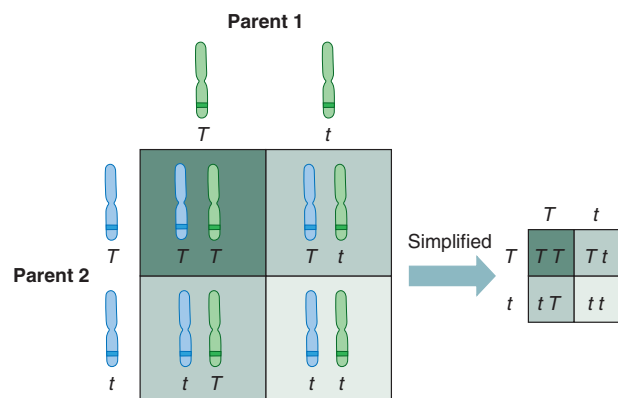


Figure 4.6 A Punnett square. A Punnett square illustrates how alleles combine in offspring. The different types of gametes of one parent are listed along the top of the square, with those of the other parent listed on the left-hand side. Each compartment displays the genotype that results when gametes that correspond to that compartment join.

square represents how particular genes in gametes join, assuming they are on different chromosomes. Experimental crosses yielded numbers of offspring that approximate these ratios.

Mendel distinguished the two genotypes resulting in tall progeny— TT from Tt —with more crosses (figure 4.7). He bred tall plants of unknown genotype with short (tt) plants. If a tall plant crossed with a tt plant produced both tall and short progeny, it was genotype Tt ; if it produced only tall plants, it must be TT .

Crossing an individual of unknown genotype with a homozygous recessive individual is called a test cross. The logic is that the homozygous recessive is the only genotype

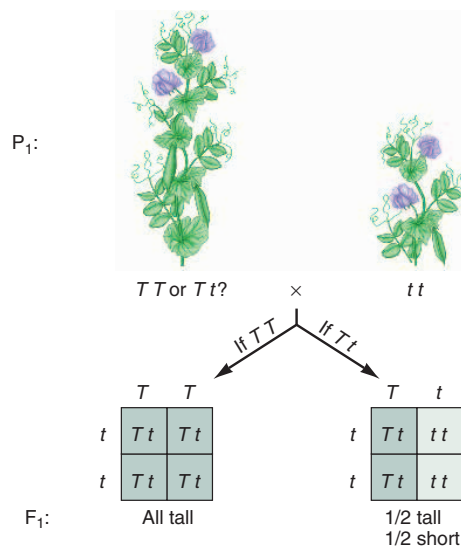


Figure 4.7 Test cross. Breeding a tall pea plant with homozygous recessive short plants reveals whether the tall plant is true-breeding (TT) or non-true-breeding (Tt). Punnett squares usually indicate only the alleles.

Table 4.1 Mendel's Law of Segregation

Experiment	Total	Dominant	Recessive	F ₂ Phenotypic Ratios
1. Seed form	7,324	5,474	1,850	2.96:1
2. Seed color	8,023	6,022	2,001	3.01:1
3. Seed coat color	929	705	224	3.15:1
4. Pod form	1,181	882	299	2.95:1
5. Pod color	580	428	152	2.82:1
6. Flower position	858	651	207	3.14:1
7. Stem length	1,064	787	277	2.84:1
				Average = 2.98:1

that can be identified by its phenotype—that is, a short plant is always *tt*. The homozygous recessive is a “known” that can reveal the unknown genotype of another individual to which it is crossed.

Key Concepts

1. Mendel deduced that “elementen” for height segregate, then combine at random with those from the opposite gamete at fertilization.
2. A homozygote has two identical alleles, and a heterozygote has two different alleles. The allele expressed in a heterozygote is dominant; the allele not expressed is recessive.
3. A monohybrid cross yields a genotypic ratio of 1:2:1 and a phenotypic ratio of 3:1.
4. Punnett squares display expected genotypic and phenotypic ratios among progeny.
5. A test cross uses a homozygous recessive individual to reveal an unknown genotype.

4.3 Single-Gene Inheritance in Humans

Mendel’s first law addresses traits determined by single genes. **Reading 4.1** describes a few unusual, single-gene traits. Inheritance of single genes is also called Mendelian, or monofactorial, inheritance. These disorders, such as sickle cell disease or muscular dystrophy, are rare compared to infectious diseases, cancer, and multifactorial disorders, affecting 1 in 10,000 or fewer individuals. Because of the rarity of single-gene diseases, getting an accurate diagnosis can be difficult because physicians are unfamiliar with the phenotype. The U.S. National Institutes of Health Undiagnosed Diseases Program is addressing this problem.

Single-gene inheritance is much more complicated than it might appear from considering such obvious traits as green or yellow pea color. Sequencing the human genome and using SNPs (points in the genome where people vary) to catalog inherited variation have revealed that the phenotypes associated with single genes are influenced by other genes as well as by environmental factors. Eye color is a good example of the new view of single-gene traits.

Eye Color

Most people have brown eyes; blue and green eyes are almost exclusively in people of European ancestry. The color of the iris is due to melanin pigments, which come in two forms—the dark brown/black eumelanin, and the red-yellow pheomelanin. In the eye, cells called melanocytes produce melanin, which is stored in structures called melanosomes in the outermost layer of the iris. People differ in the amount of melanin and number of melanosomes, but have about the same number of melanocytes in their eyes.

Nuances of eye color—light versus dark brown, clear blue versus greenish or hazel—arise from the distinctive peaks and valleys at the back of the iris. Thicker regions darken appearance of the pigments, rendering brown eyes nearly black in some parts and blue eyes closer to purple. The bluest eyes have thin irises with very little pigment. The effect of the iris surface on color is a little like the visual effect of a rough-textured canvas on paint (**figure 4.8**).

A single gene on chromosome 15, *OCA2* (MIM 611409), confers eye color by controlling melanin synthesis. If this gene is missing, albinism results, causing very pale skin and red eyes (see figure 4.17). A recessive allele of this gene confers blue color and a dominant allele confers brown. Wild type human eyes are brown because this is the most common phenotype in the global human population. But it is more complicated than this. Near the *OCA2* gene on chromosome 15 is a second gene, *HERC2*, that controls expression of the *OCA2* gene. A certain SNP in *HERC2* abolishes the control over *OCA2*, and blue eyes result. A person must inherit two copies of this SNP to have blue eyes.

If blue eye color is the disruption of a “normal” function, why has it persisted? A clue comes from evolution. The *HERC2* gene is found in many species, indicating that it is ancient. Researchers hypothesize that mutations in *HERC2* arose among long ago hunter-gatherers in Europe, and the unusual individuals with the pale eyes were, for whatever reason, more desirable as sexual partners. Over time, this sexual selection would have increased the proportion of the population with blue eyes.



Figure 4.8 Eye color. Wild type human eye color is brown; blue and green eyes stemmed from mutations or SNPs that persisted. The surface of the back of the iris contributes to the intensity of eye color.

Reading 4.1

It's All in the Genes

Do you have uncombable hair, misshapen toes or teeth, or a pigmented tongue tip? Are you unable to smell a squashed skunk, or do you sneeze repeatedly in bright sunlight? Do you lack teeth, eyebrows, eyelashes, nasal bones, thumbnails, or fingerprints? If so, your unusual trait may be one of thousands described in MIM.

Genes control whether hair is blond, brown, or black, has red highlights, and is straight, curly, or kinky. Widow's peaks, cowlicks, a whorl in the eyebrow, and white forelocks run in families; so do hairs with triangular cross-sections. Some people have multicolored hairs, like cats; others have hair in odd places, such as on the elbows, nose tip, knuckles, palms, or soles. Teeth can be missing or extra, protuberant or fused, present at birth, shovel-shaped, or "snowcapped." A person can have a grooved tongue, duckbill lips, flared ears, egg-shaped pupils, three rows of eyelashes, spotted nails, or "broad thumbs and great toes." Extra breasts are known in humans and guinea pigs, and one family's claim to genetic fame is a double nail on the littlest toe.

Unusual genetic variants can affect metabolism, producing either disease or harmless, yet noticeable, effects. Members of some families experience "urinary excretion of odoriferous component of asparagus" or "urinary excretion of beet pigment," producing a strange odor or dark pink urine after consuming the offending vegetable. In blue diaper syndrome, an infant's urine turns blue on contact with air, thanks to an inherited inability to break down the amino acid tryptophan.

One bizarre inherited illness is the Jumping Frenchmen of Maine syndrome (MIM 244100). This exaggerated startle reflex was first noted among French-Canadian lumberjacks from the Moosehead Lake area of Maine, whose ancestors were from the Beauce region of Quebec. Physicians first reported the condition at a medical conference in 1878. Geneticists videotaped the startle response in

1980, and the condition continues to appear in genetics journals. MIM offers a most vivid description:

If given a short, sudden, quick command, the affected person would respond with the appropriate action, often echoing the words of command. . . . For example, if one of them was abruptly asked to strike another, he would do so without hesitation, even if it was his mother and he had an ax in his hand.

The Jumping Frenchmen of Maine syndrome may be an extreme variant of the more common Tourette syndrome, which causes tics and other uncontrollable movements. **Figure 1** illustrates some other genetic variants.

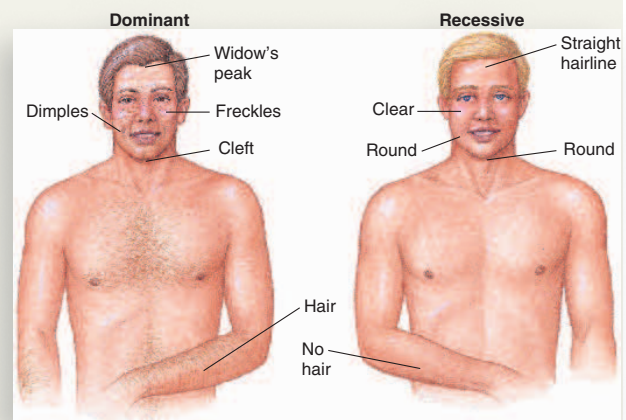


Figure 1 Inheritance of some common traits. Freckles, dimples, hairy arms, widow's peak, and a cleft chin are examples of dominant traits.

Modes of Inheritance

Despite our increasing knowledge of how genes impact each other, it is still possible, and valuable, to know who in a family has or carries a particular single-gene trait or illness. Modes of inheritance are rules that explain the common patterns of single-gene transmission. They are derived from Mendel's laws. Knowing mode of inheritance makes it possible to calculate the probability that a particular couple will have a child who inherits a particular condition. The way that a trait is passed depends on whether the gene that determines it is on an autosome or on a sex chromosome, and whether the particular allele is recessive or dominant.

In autosomal dominant inheritance, a trait can appear in either sex because an autosome carries the gene. If a child has the trait, at least one parent also has it. That is, autosomal dominant traits do not skip generations. If no offspring inherit the trait in one generation, its transmission stops because the offspring can pass on only the recessive form of the gene.

Huntington disease is an autosomal dominant condition. Henry, from section 4.1, has an affected parent and grandparent, but he has not inherited the mutation. Because his sister decided not to have children after learning she had inherited the mutation, disease transmission in their branch of the family tree stopped. The Punnett square in **figure 4.9** depicts inheritance of an autosomal dominant trait or condition, and **table 4.2** summarizes the criteria. Many autosomal dominant diseases do not cause symptoms until adulthood. The *Bioethics: Choices for the Future* box on page 78 looks at the dilemma that adult-onset inherited diseases presents for reproductive choices.

An autosomal recessive trait can appear in either sex. Affected individuals have a homozygous recessive genotype, whereas in heterozygotes (carriers) the wild type allele masks expression of the mutant allele. Ellen, the little girl with CF described in section 4.1, inherited a mutant allele from each of her carrier parents. They were unaffected because they each

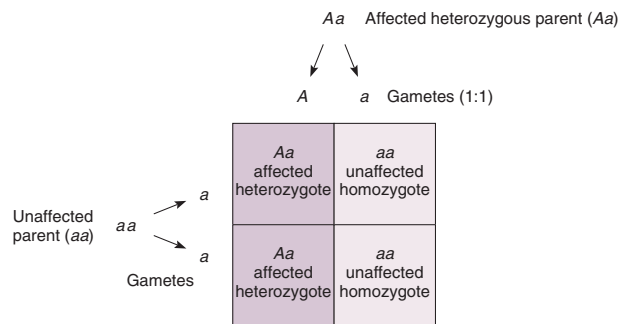


Figure 4.9 Autosomal dominant inheritance. When one parent has an autosomal dominant condition and the other does not, each offspring has a 50 percent probability of inheriting the mutant allele and the condition. The affected parent is Aa here, and not AA , because for many dominant disorders, the homozygous dominant (AA) phenotype is either lethal or very rare because both parents of the person with the AA genotype would have to have the disorder.

also had a dominant allele that encodes enough functional protein for health. **Figure 4.10** depicts the autosomal recessive inheritance pattern of a harmless trait, curly hair.

Mendel's first law can be used to calculate the probability that an individual will have either of two phenotypes. The probabilities of each possible genotype are added. For example, the chance that a child whose parents are both carriers of cystic fibrosis will *not* have the condition is the sum of the probability that she has inherited two normal alleles ($1/4$) plus the chance that she herself is a heterozygote ($1/2$), or $3/4$. Note that this also equals 1 minus the probability that she is homozygous recessive and has the condition.

The ratios that Mendel's first law predicts for autosomal recessive inheritance apply to each offspring anew. If a couple has a child with an autosomal recessive illness, each of their next children faces the same 25 percent risk of inheriting the condition.

Most autosomal recessive conditions appear unexpectedly in families. However, a situation in which an autosomal recessive condition is more likely to recur is when blood relatives have children together. The higher risk of having a child with a particular autosomal recessive condition is because the related parents may carry the same alleles inherited from an ancestor that they have in common, such as a grandparent.

Table 4.2 Criteria for an Autosomal Dominant Trait

1. Males and females can be affected. Male-to-male transmission can occur.
2. Males and females transmit the trait with equal frequency.
3. Successive generations are affected.
4. Transmission stops after a generation in which no one is affected.

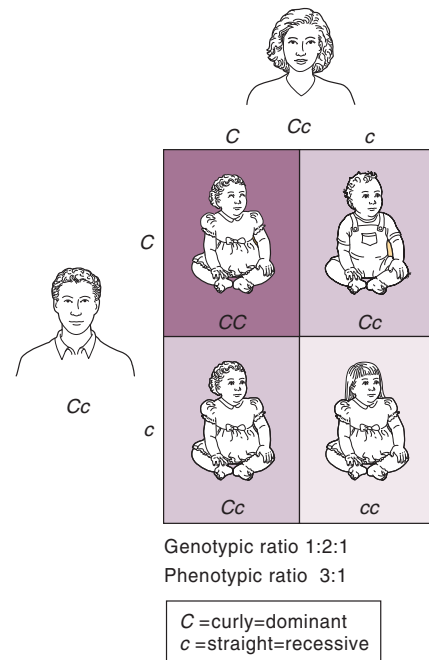


Figure 4.10 Autosomal recessive inheritance. A 1:2:1 genotypic ratio results from a monohybrid cross, whether in peas or people. Curly hair (C) is dominant to straight hair (c). This pedigree depicts a monohybrid cross for curly hair.

Marriage between relatives introduces **consanguinity**, which means “shared blood”—a figurative description, since genes are not passed in blood.

Logic explains why consanguinity raises risk of inheriting autosomal recessive diseases. An unrelated man and woman have eight different grandparents, but first cousins have only six, because they share one pair through their parents, who are siblings (see figure 4.16c). That is, the probability of two relatives inheriting the same disease-causing recessive allele is greater than that of two unrelated people having the same allele by chance.

The nature of the phenotype is important when evaluating the transmission of single-gene traits. Some diseases are too severe for people to live long enough or feel well enough to have children. For example, each adult sibling of a person who is a known carrier of Tay-Sachs disease has a two-thirds chance of being a carrier. The probability is two-thirds, and not one-half, because there are only two genotypes possible for an adult—homozygous for the wild type allele or a carrier who inherits the mutant allele from either mother or father. A homozygous recessive individual would not have survived childhood, due to brain degeneration.

Geneticists who study human traits and illnesses can hardly set up crosses as Mendel did, but they can pool information from families whose members have the same trait or illness. Consider a simplified example of 50 couples in whom both partners are carriers of sickle cell disease. If 100 children are born, about 25 of them would be expected to have sickle cell disease. Of the remaining 75, theoretically 50 would be

Bioethics: Choices for the Future

When Diagnosing a Fetus Also Diagnoses a Parent: Huntington Disease (HD)

When Peter and Martha were 24 and expecting their first child, they learned that Peter's mother, who was adopted, had early signs of HD. Her clumsiness and slurred speech would eventually progress to a near-constant writhing and repetitive, dance-like movements, until she would die, probably of infection, in 15 to 20 years.

A genetic counselor told the couple that because HD is autosomal dominant, Peter had a 50 percent (1 in 2) chance of having inherited the condition. He was considered "at-risk" until (and if) symptoms began. However, he could take a "predictive" genetic test that would reveal whether he had inherited HD. If he had, he would be considered "pre-manifest" before signs and symptoms begin. The mutation extends the wild type gene and consists of extra repeats of a three-base DNA sequence. The number of extra repeats could be used to predict the time until symptom onset—but Peter didn't want that information. Martha, however, felt differently. She did not want to have a child who would have HD. Could the fetus be checked for the mutation? Then, if the disease had been inherited, the couple could end the pregnancy.

The problem was that if the fetus had the mutation, then so did Peter, and he didn't want to know. He also questioned ending a pregnancy, considering that symptoms do not begin until later in life, by which time treatment might exist.

The couple researched how others had handled the dilemmas that predictive testing for an untreatable, adult-onset illness raise. They read that only about 10 percent of people offered the test for HD take it, and those that do can handle the results well. After much soul-searching, Martha chose to respect Peter's wishes. He and the fetus were not tested. So far, both are healthy.

Postscript: Although Peter still does not wish to know whether he will get HD, he is participating in a large-scale study. He and 1,000 other at-risk individuals have given a blood sample to researchers. The participants will be followed medically for 10 years to identify the very earliest signs and symptoms of the disease—which might suggest treatments, perhaps even existing drugs.

Questions for Discussion

1. Some geneticists advise that people under 18 not be tested for the HD mutation, unless they are very mature and well-informed. Do you agree with this practice?
2. Choose an autosomal dominant condition from MIM, familyvillage.org, genetests.org, or wikipedia, and argue for or against testing children for the mutation.
3. Discuss issues of privacy that may arise in genetic counseling for an autosomal dominant condition.

carriers like their parents, and the remaining 25 would have two wild type alleles. **Table 4.3** lists criteria for an autosomal recessive trait.

Table 4.3 Criteria for an Autosomal Recessive Trait

1. Males and females can be affected.
2. Affected males and females can transmit the gene, unless it causes death before reproductive age.
3. The trait can skip generations.
4. Parents of an affected individual are heterozygous or have the trait.

+ Solving a Problem

Segregation

Using Mendel's laws to predict phenotypes and genotypes requires a careful reading of the problem to identify and organize relevant information. Sometimes common sense is useful, too. The following general steps can help to solve a problem that addresses Mendel's first law, which describes the inheritance of a single-gene trait.

1. List all possible genotypes and phenotypes for the trait.
2. Determine the genotypes of the individuals in the first (P_1) generation. Use information about those individuals' parents.
3. After deducing genotypes, derive the possible alleles in gametes each individual produces.
4. Unite these gametes in all combinations to reveal all possible genotypes. Calculate ratios for the F_1 generation.
5. To extend predictions to the F_2 generation, use the genotypes of the specified F_1 individuals and repeat steps 3 and 4.

As an example, consider curly hair, depicted in figure 4.10. If C is the dominant allele, conferring curliness, and c is the recessive allele, then both CC and Cc genotypes result in curly hair. A person with a cc genotype has the straight hair phenotype.

Wendy has beautiful curls, and her husband Rick has straight hair. Wendy's father is bald, but once had curly hair, and her mother has stick-straight hair. What is the probability that Wendy and Rick's child will have straight hair? Steps 1 through 5 solve the problem:

1. State possible genotypes: CC , Cc = curly; cc = straight
2. Determine genotypes: Rick must be cc , because his hair is straight. Wendy must be Cc , because her mother has straight hair and therefore gave her a c allele.

3. Determine gametes: Rick's sperm carry only *c*. Half of Wendy's oocytes carry *C*, and half carry *c*.

4. Unite the gametes: Wendy

		<i>C</i>	<i>c</i>
Rick	<i>c</i>	<i>Cc</i>	<i>cc</i>
	<i>c</i>	<i>Cc</i>	<i>cc</i>

5. Conclusion: Each child of Wendy and Rick has a 50 percent chance of having curly hair (*Cc*) and a 50 percent chance of having straight hair (*cc*).

On the Meaning of Dominance and Recessiveness

Determining whether an allele is dominant or recessive is critical in determining risk of inheriting a particular condition (phenotype). Dominance and recessiveness arise from the genotype, and reflect the characteristics or abundance of a protein.

Mendel based his definitions of dominance and recessiveness on what he could see—one allele masking the other. Today we can often add a cellular or molecular explanation. Consider inborn errors of metabolism caused by absent enzymes. These disorders are typically recessive because the half normal amount of the enzyme that a carrier produces is usually sufficient to maintain health. The one normal allele, therefore, compensates for the mutant one, to which it is dominant. The situation is similar in pea plants. Short stem length results from deficiency of an enzyme that activates a growth hormone, but the *Tt* plants produce enough hormone to attain the same height as *TT* plants.

A recessive trait is said to arise from a “loss of function” because the recessive allele usually causes the loss of normal protein production and function. In contrast, some dominantly inherited disorders are said to be due to a “gain of function,” because they result from the action of an abnormal protein that interferes with the function of the normal protein. Huntington disease is a “gain of function” disorder. In HD, the dominant mutant allele encodes an abnormally long protein that prevents the normal protein from functioning in certain brain cells. Huntington disease is a gain of function because individuals who are missing one copy of the gene do not have the illness. That is, the protein encoded by the mutant HD allele must be abnormal, not absent, to cause the disease.

Recessive disorders tend to be more severe, and produce symptoms earlier than dominant disorders. Disease-causing recessive alleles remain in populations because healthy heterozygotes pass them to future generations. In contrast, if a dominant mutation arises that harms health early in life, people who have the allele are either too ill or do not survive long enough to reproduce. The allele eventually becomes rare in the population unless it arises anew by mutation. Dominant disorders whose symptoms do not appear until adulthood, or that do not drastically disrupt health, remain in a population because they do not affect health until after a person has reproduced.

Key Concepts

1. Modes of inheritance reveal whether a single-gene trait is dominant or recessive and whether the gene that controls it is carried on an autosome or a sex chromosome.
2. Autosomal dominant traits do not skip generations and can affect both sexes; autosomal recessive traits can skip generations and can affect both sexes.
3. Rare autosomal recessive disorders sometimes recur in families when parents are related.
4. Mendel's first law, which can predict the probability that a child will inherit a single-gene trait, applies anew to each child.
5. Genetic problems are solved with logic and by applying Mendel's laws to follow gametes.
6. Dominance is the ability of a protein encoded by one allele to compensate for a missing or abnormal protein encoded by another allele.

4.4 Following the Inheritance of Two Genes—Independent Assortment

The law of segregation follows the inheritance of two alleles for a single gene. In a second set of experiments, Mendel examined the inheritance of two different traits, each attributable to a gene with two different alleles.

Mendel's Second Law

The second law, the **law of independent assortment**, states that for two genes on different chromosomes, the inheritance of one does not influence the chance of inheriting the other. The two genes thus “independently assort” because they are packaged into gametes at random (**figure 4.11**). Two genes that are far apart on the same chromosome also appear to independently assort, because so many crossovers occur between them that it is as if they are carried on separate chromosomes (see **figure 3.5**).

Mendel looked at seed shape, which was either round or wrinkled (determined by the *R* gene), and seed color, which was either yellow or green (determined by the *Y* gene). When he crossed true-breeding plants that had round, yellow seeds to true-breeding plants that had wrinkled, green seeds, all the progeny had round, yellow seeds. These offspring were double heterozygotes, or dihybrids, of genotype *RrYy*. From their appearance, Mendel deduced that round is dominant to wrinkled, and yellow to green.

Next, he self-crossed the dihybrid plants in a **dihybrid cross**, so named because two genes and traits are followed. Mendel found four types of seeds in the next, third generation: 315 plants with round, yellow seeds; 108 plants with round,

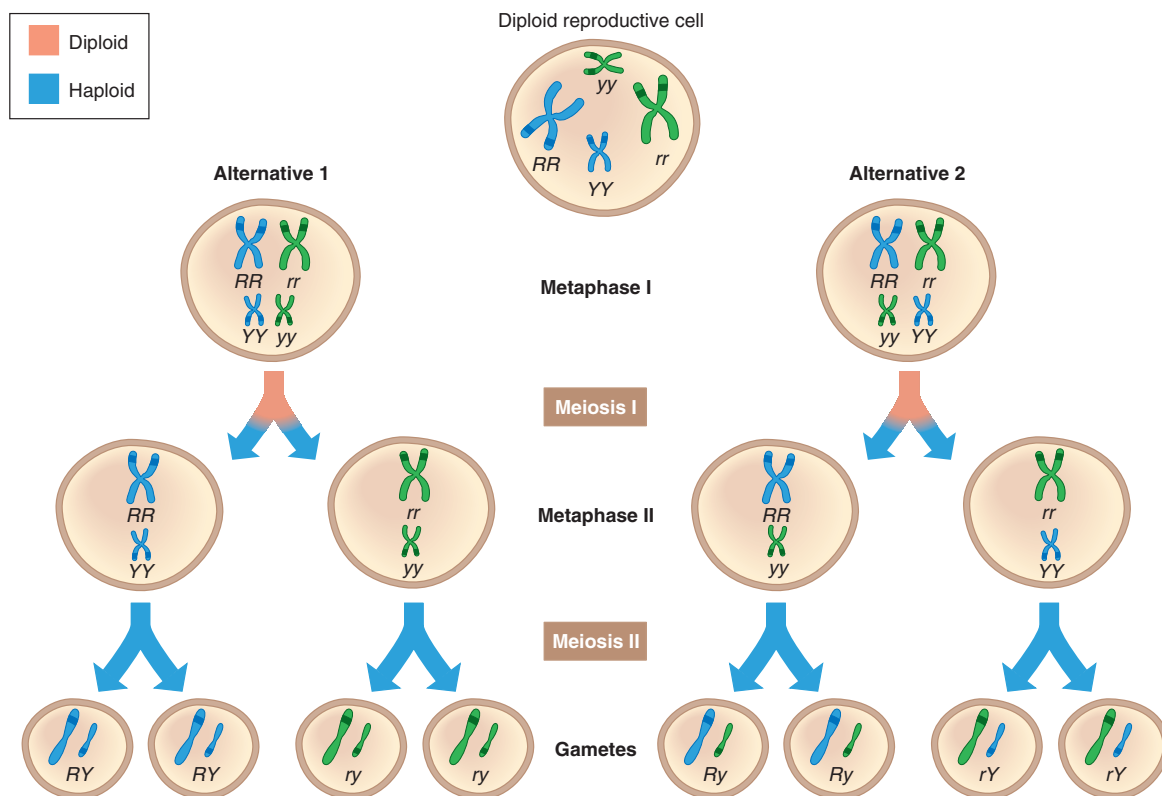


Figure 4.11 Mendel's second law—-independent assortment. The independent assortment of genes carried on different chromosomes results from the random alignment of chromosome pairs during metaphase of meiosis I. An individual of genotype $RrYy$, for example, manufactures four types of gametes, containing the dominant alleles of both genes (RY), the recessive alleles of both genes (ry), and a dominant allele of one with a recessive allele of the other (Ry or rY). The allele combination depends upon which chromosomes are packaged together in a gamete—and this happens at random.

green seeds; 101 plants with wrinkled, yellow seeds; and 32 plants with wrinkled, green seeds. These classes occurred in a ratio of 9:3:3:1.

Mendel then took each plant from the third generation and crossed it to plants with wrinkled, green seeds (genotype $rryy$). These test crosses established whether each plant in the third generation was true-breeding for both genes (genotypes $RRYY$ or $rryy$), true-breeding for one gene but heterozygous for the other (genotypes $RRYy$, $RrYY$, $rRYy$, or $Rryy$), or heterozygous for both genes (genotype $RrYy$). Mendel could explain the 9:3:3:1 proportion of progeny classes only if one gene does not influence transmission of the other. Each parent would produce equal numbers of four different types of gametes: RY , Ry , rY , and ry . Note that each of these combinations has one gene for each trait. A Punnett square for this cross shows that the four types of seeds:

1. round, yellow ($RRYY$, $RrYY$, $RRYy$, and $RrYy$)
2. round, green ($RRyy$ and $Rryy$)
3. wrinkled, yellow ($rrYY$ and $rRYy$) and
4. wrinkled, green ($rryy$)

are present in the ratio 9:3:3:1, just as Mendel found (figure 4.12).

+ Solving a Problem

Following More Than One Segregating Gene

A Punnett square for three genes has 64 boxes; for four genes, 256 boxes. An easier way to predict genotypes and phenotypes in multi-gene crosses is to use the mathematical laws of probability on which Punnett squares are based. Probability predicts the likelihood of an event.

An application of probability theory called the product rule can predict the chance that parents with known genotypes can produce offspring of a particular genotype. The product rule states that the chance that two independent events will both occur equals the product of the chance that either event will occur alone. Consider the probability of obtaining a plant with wrinkled, green peas (genotype $rryy$) from dihybrid ($RrYy$) parents. Do the reasoning for one gene at a time, then multiply the results (figure 4.13).

A Punnett square for Rr crossed to Rr shows that the probability of Rr plants producing rr progeny is 25 percent, or $1/4$. Similarly, the chance of two Yy plants producing a yy plant is $1/4$. Therefore, the chance of dihybrid parents ($RrYy$) producing homozygous recessive ($rryy$) offspring is $1/4$ multiplied by $1/4$, or $1/16$. Now consult the 16-box Punnett square for Mendel's

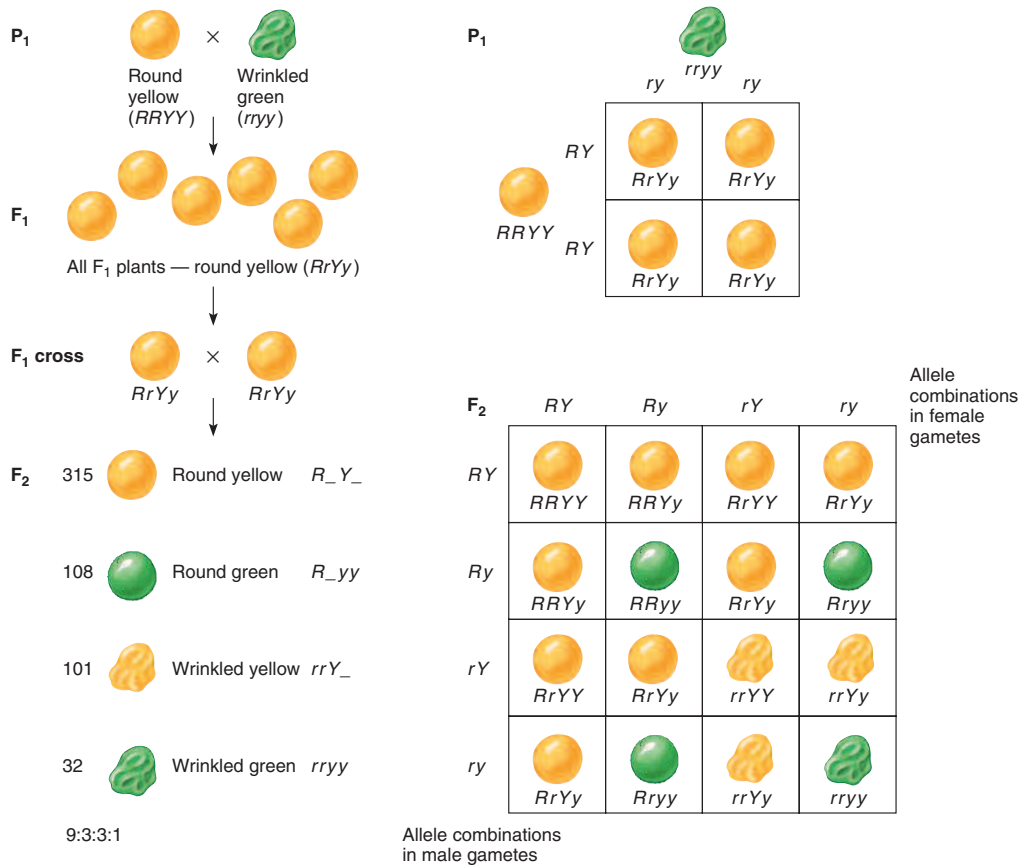


Figure 4.12 Plotting a dihybrid cross. A Punnett square can represent the random combinations of gametes produced by dihybrid individuals. An underline in a genotype (in the F₂ generation) indicates that either a dominant or a recessive allele is possible. The numbers in the F₂ generation are Mendel's experimental data.

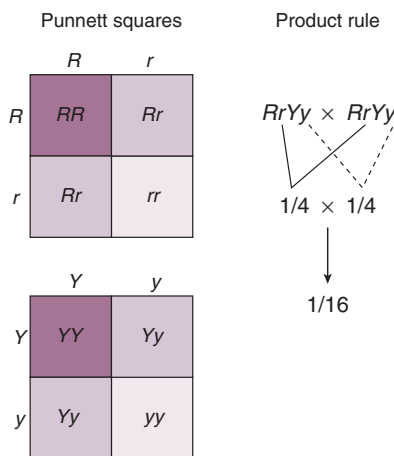


Figure 4.13 The product rule.

dihybrid cross again (figure 4.12). Only one of the 16 boxes is rryy, just as the product rule predicts. **Figure 4.14** shows how probability and Punnett squares can be used to predict offspring genotypes and phenotypes for three human traits simultaneously.

Until recently, Mendel's second law has not been as useful in medical genetics as the first law, because not enough genes were identified to follow the transmission of two or more traits at a time. It is common now to screen for hundreds or thousands of alleles or expressed genes at once. Computer analysis of many gene combinations has largely replaced Punnett squares.

Key Concepts

1. Mendel's law of independent assortment considers genes transmitted on different chromosomes.
2. In a dihybrid cross of heterozygotes for seed color and shape, the phenotypic ratio of 9:3:3:1 revealed that transmission of one gene does not influence that of another.
3. Meiotic events explain independent assortment.
4. Punnett squares and probability can be used to follow independent assortment.
5. Today, computers analyze many genes at a time.

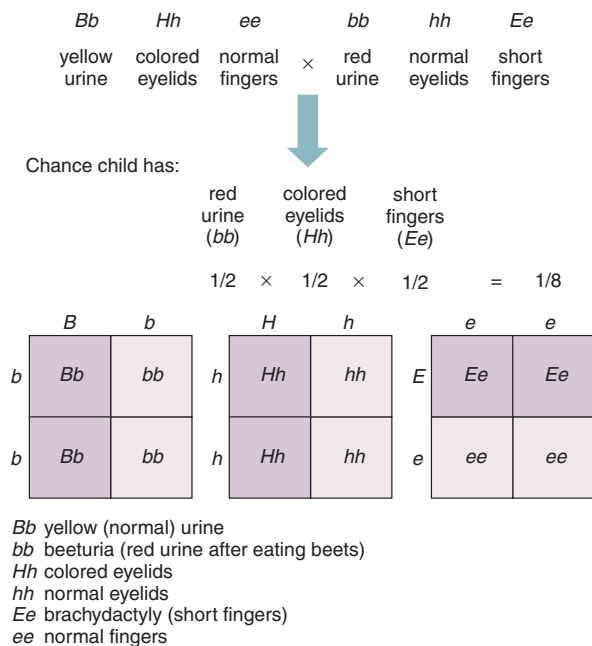


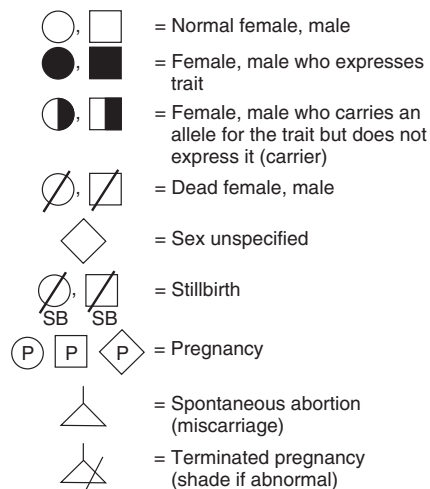
Figure 4.14 Using probability to track three traits. A man with normal urine, colored eyelids, and normal fingers wants to have children with a woman who has red urine after she eats beets, normal eyelids, and short fingers. The chance that a child of theirs will have red urine after eating beets, colored eyelids, and short fingers is $1/8$.

4.5 Pedigree Analysis

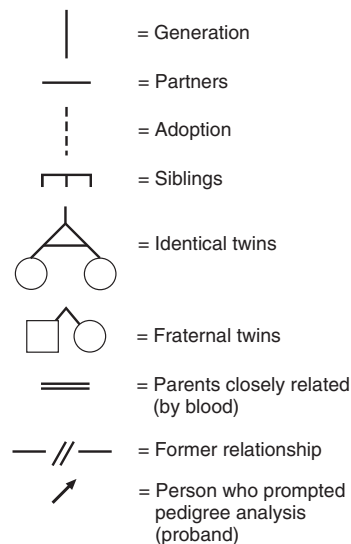
For researchers, families are tools, and the bigger the family the better—the more children in a generation, the easier it is to discern modes of inheritance. Charts called **pedigrees** display family relationships and depict which relatives have specific phenotypes and, sometimes, genotypes. A human pedigree serves the same purpose as one for purebred dogs or cats or thoroughbred horses—it represents relationships. A pedigree in genetics differs from a family tree in genealogy, and from a genogram in social work, in that it indicates disorders or traits as well as relationships and ancestry. Pedigrees may also include molecular data, test results, haplotypes (genes or SNPs linked in segments on a chromosome), and even genome-wide association study information.

A pedigree consists of lines that connect shapes. Vertical lines represent generations; horizontal lines that connect two shapes at their centers depict partners; shapes connected by vertical lines that are joined horizontally represent siblings. Squares indicate males; circles, females; and diamonds, individuals of unspecified sex. Roman numerals designate generations. Arabic numerals or names indicate individuals. **Figure 4.15** shows these and other commonly used pedigree symbols. Colored or shaded shapes indicate individuals who express a trait, and half-filled shapes are known carriers. A genetic counselor may sketch a pedigree while interviewing a client, then use a computer program and add test results that indicate genotypes.

Symbols



Lines



Numbers

Roman numerals = generations

Arabic numerals = individuals in a generation

Figure 4.15 Pedigree components. Symbols representing individuals are connected to form pedigree charts, which display the inheritance patterns of particular traits.

Pedigrees Then and Now

The earliest pedigrees were strictly genealogical, not indicating traits. **Figure 4.16** shows such a pedigree for a highly inbred part of the ancient Egyptian royal family. The term *pedigree* arose in the fifteenth century, from the French *pie de grue*, which means “crane’s foot.” Pedigrees at that time, typically depicting large

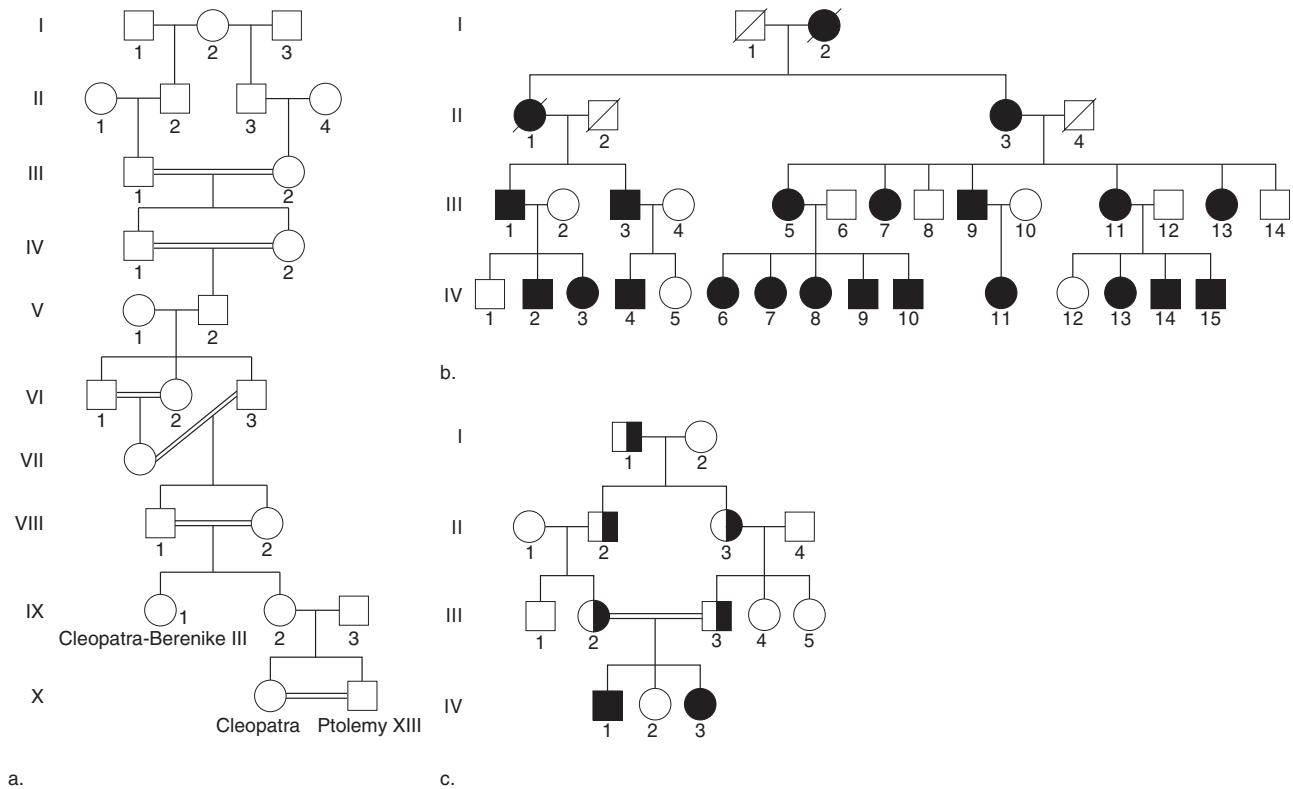


Figure 4.16 Unusual pedigrees. (a) A partial pedigree of Egypt’s Ptolemy dynasty shows only genealogy, not traits. It appears almost ladderlike because of the extensive inbreeding. From 323 b.c. to Cleopatra’s death in 30 b.c., the family experienced one pairing between cousins related through half-brothers (generation III), four brother-sister pairings (generations IV, VI, VIII, and X), and an uncle-niece relationship (generations VI and VII). Cleopatra married her brother, Ptolemy XIII, when he was 10 years old! These marriage patterns were an attempt to preserve the royal blood. (b) In contrast to the Egyptian pedigree, a family with polydactyly (extra fingers and toes) extends laterally, with many children and depiction of the family’s trait in the filled-in symbols. (c) This pedigree shows marriage of first cousins. They share one set of grandparents, and therefore risk passing on the same recessive alleles to offspring.

families, showed parents linked by curved lines to their many offspring. The overall diagram often resembled a bird’s foot.

One of the first pedigrees to trace an inherited illness was an extensive family tree of several European royal families, indicating which members had the clotting disorder hemophilia (see figure 6.8). The mutant gene probably originated in Queen Victoria of England in the nineteenth century. In 1845, a genealogist named Pliny Earle constructed a pedigree of a family with colorblindness using musical notation—half notes for unaffected females, quarter notes for colorblind females, and filled-in and squared-off notes to represent the many colorblind males. In the early twentieth century, eugenicists tried to use pedigrees to show that traits such as criminality, feeble-mindedness, and promiscuity were the consequence of faulty genes. Figure 15.16 shows one of these pedigrees.

Today, pedigrees are important both for helping families identify the risk of transmitting an inherited illness and as starting points for identifying a gene from the human genome sequence. People who have kept meticulous family records are invaluable in helping researchers follow the inheritance of particular genes in groups such as the Mormons and the Amish. Very

large pedigrees can provide information on many individuals with a particular rare disorder. The researchers can then search affected individuals’ DNA to identify a particular sequence they have all inherited that is not found in healthy family members. This is where the causative mutation lies. Discovery of the gene that causes HD, for example, took researchers to a remote village in Venezuela to study an enormous family. The gene was eventually traced to a Portuguese sailor who introduced the mutation in the nineteenth century (Reading 22.1).

Pedigrees Display Mendel’s Laws

Visual learners can easily “see” a mode of inheritance in a pedigree. Consider a pedigree for an autosomal recessive trait, albinism. Homozygous recessive individuals in the third (F_2) generation lack an enzyme necessary to manufacture the pigment melanin and, as a result, hair and skin are very pale (figure 4.17) Their parents are inferred to be heterozygotes (carriers). One partner from each pair of grandparents must also be a carrier. For some disorders, carriers can be identified by detecting half the wild type amount of a key biochemical.

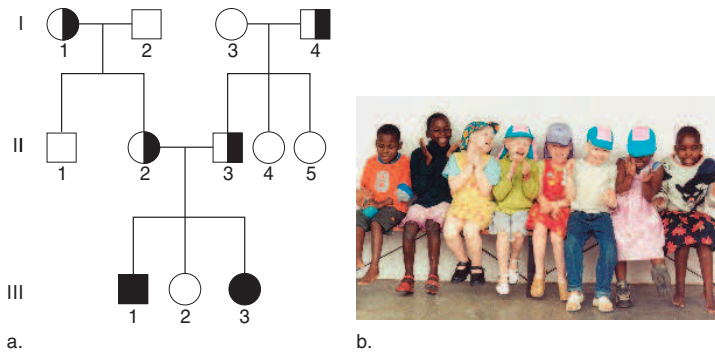


Figure 4.17 Albinism is autosomal recessive. (a) Albinism affects males and females and can skip generations, as it does here in generations I and II in this pedigree. Homozygous recessive individuals lack an enzyme needed to produce melanin, which colors the eyes, skin, and hair. The South African children with albinism in (b) stand out from their dark-skinned classmates.

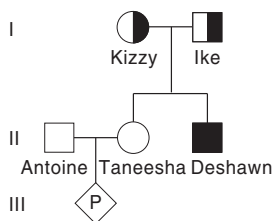
An autosomal dominant trait does not skip generations and can affect both sexes. A typical pedigree for an autosomal dominant trait has some squares and circles filled in to indicate affected individuals in each generation. The inset in figure 4.1 and figure 4.16b illustrate autosomal dominant inheritance.

A pedigree may be inconclusive, which means that either autosomal recessive or autosomal dominant inheritance can explain the pattern of filled-in symbols. **Figure 4.18** shows one such pedigree, for a type of hair loss called alopecia areata (MIM 104000). According to the pedigree, this trait can be passed in an autosomal dominant mode because it affects both males and females and is present in every generation. However, the pedigree can also depict autosomal recessive inheritance if the individuals represented by unfilled symbols are carriers. Inconclusive pedigrees tend to arise when families are small and the trait is not severe enough to impair fertility.

⊕ Solving a Problem

Conditional Probability

Genetic counselors are asked to predict the probability that a condition will occur in a particular individual. Mendel's laws, pedigrees, and Punnett squares provide clues, as do logic



a. Taneesha's brother, Deshawn, has sickle cell disease.

		Kizzy	
		S	s
Ike	S	SS	Ss
	s	Ss	ss

b. Probability that Taneesha is a carrier: $\frac{2}{3}$

		Taneesha	
		S	s
Antoine	S	SS	Ss
	s	SS	Ss

c. If Taneesha is a carrier, chance that fetus is a carrier: $\frac{1}{2}$

$$\text{Total probability} = \frac{2}{3} \times \frac{1}{2} = \frac{1}{3}$$

Figure 4.19 Making predictions. Taneesha's brother, Deshawn, has sickle cell disease (a). Taneesha wonders if her fetus has inherited the sickle cell allele from her. First, she must calculate the chance that she is a carrier. The Punnett square in (b) shows that this risk is 2 in 3. (She must be genotype SS or Ss, but cannot be ss because she does not have the disease.) The risk that the fetus is a carrier, assuming that the father is not a carrier, is half Taneesha's risk of being a carrier, or 1 in 3 (c).

Figure 4.18 An inconclusive pedigree. This pedigree could account for an autosomal dominant trait or an autosomal recessive trait that does not prevent affected individuals from having children. (Unfilled symbols could represent carriers.)

and common sense. In the family depicted in **figure 4.19**, Deshawn has sickle cell disease, which is autosomal recessive. His unaffected parents, Kizzy and Ike, must each be heterozygotes. Deshawn's sister, Taneesha, also healthy, is expecting her first child. Taneesha's husband, Antoine, has no family history of sickle cell disease. Taneesha wants to know the risk that her child will inherit the mutant allele from her and be a carrier.

Taneesha raises two questions. First, what is the risk that she is a carrier? Because Taneesha is the product of a monohybrid cross, and she is not homozygous recessive, she has a 2 in 3 chance of being a carrier. If so, the chance that she will transmit the mutant allele is 1 in 2, because she has two copies of the gene, and only one allele goes into each gamete. To calculate the overall risk to Taneesha's child, multiply the probability that she is a carrier by the chance that, if she is, she will transmit the mutant allele. This result, following two events, is a conditional probability, because the likelihood of the second event—the child being a carrier—depends upon the first event—that Taneesha is a carrier. If we assume Antoine is not a carrier, Taneesha's chance of having a child who carries the mutant allele is $\frac{2}{3}$ times $\frac{1}{2}$, which equals $\frac{2}{6}$, or $\frac{1}{3}$.

Pedigrees may be difficult to construct or interpret for several reasons. People sometimes hesitate to supply information because they are embarrassed by symptoms affecting behavior or mental stability. Family relationships can be complicated by adoption, children born out of wedlock, serial relationships, blended families, and assisted reproductive technologies (see chapter 21). Many people cannot trace their families back far enough to reveal a mode of inheritance.

Reading 4.2 is a closer look at cystic fibrosis, which is one of the most common single-gene disorders.

Key Concepts

1. Pedigrees depict family relationships and the transmission of inherited traits. Squares represent males, and circles, females; horizontal lines link partners, vertical lines show generations, and elevated horizontal lines depict siblings. Heterozygote symbols are half-shaded, and symbols for individuals who express a trait are completely shaded.
2. Pedigrees can reveal modes of inheritance. They and Punnett squares apply Mendel's laws to predict the recurrence risks of inherited disorders or traits.

Reading 4.2

Cystic Fibrosis, Then and Now

Alex Deford was born in 1972, daughter of sportswriter Frank Deford. Alex did not gain much weight as an infant, and as a young child suffered severe lung infections and digestive difficulties before dying of CF at age 8. Wrote her father about his feelings at the time of diagnosis in his book, *Alex, The Life of a Child*:

"I went to the encyclopedia and read about this cystic fibrosis. To me, at that point, it was one of those vague diseases you hear about now and then . . . One out of every 20 whites carries the defective gene, as I do, as Carol does, as perhaps 10 million other Americans do."

In the 1970s, it was common for a child with CF not to survive to see adolescence. Today, thanks to treatments and therapies to minimize symptoms, people with CF are living much longer. Within a decade, more than half of all people with the disease will be over the age of 18, and some are living normal life expectancies. Identification of more than 1,600 mutations has revealed milder guises of the condition, such as chronic sinus infections and/or bronchitis.

Diagnosis

CF is caused by faulty chloride ion channels in cells lining passageways, such as in the lungs and pancreas. The result is thickened secretions that impair breathing and digestion of fats.

Diagnosis of CF is based on any one of the following phenotypes:

- Sinus and lung disease—The bronchial tubes may be dilated, inflamed, and collapsed, and nasal polyps and infected sinuses are common. Pneumonia is caused by specific pathogens.
- Pancreatic insufficiency causes large, fatty bowel movements, malnutrition from poor fat absorption, and intestinal blockage.
- Males lack the ductus deferens, blocking sperm.

The course of CF varies greatly, but in general people lose from 1 to 4 percent of lung function a year. It may be episodic, with periods

of fatigue, shortness of breath, a productive cough, weight loss, and poor lung function.

People with CF are highly susceptible to pneumonia caused by a few types of bacteria. Children are infected with *Staphylococcus aureus* and *Hemophilus influenzae*, and adults with *Pseudomonas aeruginosa* and *Burkholderia cepacia*. Particularly dangerous is *B. cepacia*, which not only infects rapidly, but easily spreads among patients, such as children at summer camp. However, these bacterial infections may seem predominant only because we can grow the microbes in the lab. When researchers sequenced bacterial DNA in lung fluid from children with CF, they discovered genes representing more than 60 species of bacteria! It's clear that we have more to learn about the role of infection in CF.

Treatment

A combination of treatments has significantly prolonged and improved the lives of many with CF. An acute attack is treated with 2 to 3 weeks of intravenous delivery of two antibiotics, plus airway clearance therapy three to four times a day. Daily therapy to treat chronic symptoms takes an hour or two and combats obstruction, infection, and inflammation in the lungs. It includes:

- Airway clearance from manual exercises or wearing a vibrating vest.
- An inhaled drug consisting of a DNA-digesting enzyme that destroys white blood cells at infection sites, loosening sputum.
- Saline solution inhaled several times a day.
- Inhaled antibiotics.
- Anti-inflammatory drugs.

In some patients, the part of the pancreas that functions as an endocrine gland becomes damaged, leading to CF-related diabetes. These patients take short-acting insulin with meals. This type of

(continued)

(continued)

diabetes cannot be well controlled with diet because the patients need to consume very-high-caloric meals to compensate for their impaired fat digestion.

Each year about 150 people with CF receive double lung transplants. About half of them survive 5 years. Patients who benefit most from a transplant are those with lung function less than 30 percent of normal who are frequently hospitalized and have great difficulty with everyday activities.

On a more philosophical note, progress in CF presents powerful evidence against genetic determinism—that is, there is much we can do to combat an inherited phenotype. The median survival time today is close to 40 years—we've come a long way since Alex Deford lost her battle with CF (figure 1).

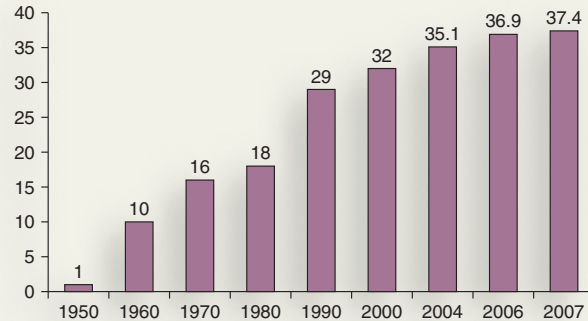


Figure 1 Living with cystic fibrosis. Many people who have CF are surviving to adulthood.

Summary

4.1 A Tale of Two Families

1. **Modes of inheritance** are the patterns in which single-gene traits and disorders occur in families.

4.2 Following the Inheritance of One Gene—Segregation

2. Mendel's laws, based on pea plant crosses, derive from the actions of chromosomes during meiosis. They apply to all diploid organisms.
3. Mendel used statistics to investigate why some traits vanish in hybrids. The **law of segregation** states that alleles of a gene are distributed into separate gametes during meiosis. Mendel demonstrated this in seven traits in pea plants.
4. A diploid individual with two identical alleles of a gene is **homozygous**. A **heterozygote** has two different alleles of a gene. A gene may have many alleles.
5. A **dominant** allele masks the expression of a **recessive** allele. An individual may be homozygous dominant, homozygous recessive, or heterozygous.
6. When Mendel crossed two true-breeding types, then bred the resulting hybrids to each other, the two variants of the trait appeared in a 3:1 phenotypic ratio. Crossing these progeny further revealed a genotypic ratio of 1:2:1.
7. A **Punnett square** follows the transmission of alleles and is based on probability.

4.3 Single-Gene Inheritance in Humans

8. Eye color illustrates how a single-gene trait can be affected by other genes.

9. In **autosomal dominant** inheritance, males and females may be affected, and the trait does not skip generations.
10. Inheritance of an **autosomal recessive** trait may affect either males or females and may skip generations. Autosomal recessive conditions are more likely to occur in families with **consanguinity**. Recessive disorders tend to be more severe and cause symptoms earlier than dominant disorders.
11. Genetic problems can be solved by tracing alleles as gametes form and then combine in a new individual.
12. Dominance and recessiveness reflect how alleles affect the abundance or activity of the gene's protein product.

4.4 Following the Inheritance of Two Genes—Independent Assortment

13. Mendel's second law, the **law of independent assortment**, follows the transmission of two or more genes on different chromosomes. A random assortment of maternally and paternally derived chromosomes during meiosis results in gametes that have different combinations of these genes.
14. The chance that two independent genetic events will both occur is equal to the product of the probabilities that each event will occur. This product rule is useful in calculating the risk that individuals will inherit a particular genotype and in following the inheritance of genes on different chromosomes.

4.5 Pedigree Analysis

15. A **pedigree** is a chart that depicts family relationships and patterns of inheritance for particular traits. A pedigree can be inconclusive.

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Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

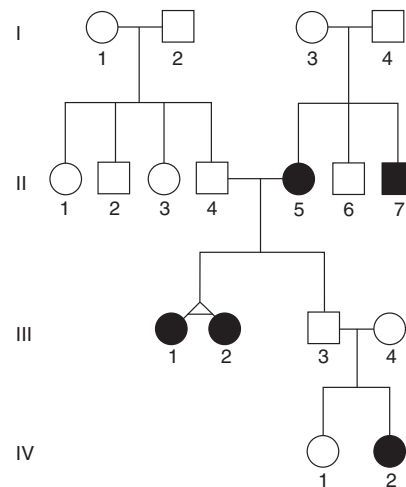
- How does meiosis explain Mendel's laws?
- How was Mendel able to derive the two laws of inheritance without knowing about chromosomes?
- Distinguish between
 - autosomal recessive and autosomal dominant inheritance.
 - Mendel's first and second laws.
 - a homozygote and a heterozygote.
 - a monohybrid and a dihybrid cross.
 - a Punnett square and a pedigree.
- Why would Mendel's results for the dihybrid cross have been different if the genes for the traits he followed were located near each other on the same chromosome?
- Why are extremely rare autosomal recessive disorders more likely to appear in families in which blood relatives have children together?
- How does the pedigree of the ancient Egyptian royal family in figure 4.16a differ from a pedigree a genetic counselor might use today?
- People who have Huntington disease inherit one mutant and one normal allele. How would an individual homozygous dominant for the condition be conceived?
- What is the probability that two individuals with an autosomal recessive trait, such as albinism, will have a child with the same genotype and phenotype as they have?

Applied Questions

- Predict the phenotypic and genotypic ratios for crossing the following pea plants:
 - short \times short
 - short \times true-breeding tall
 - true-breeding tall \times true-breeding tall
- What are the genotypes of the pea plants that would have to be bred to yield one plant with restricted pods for every three plants with inflated pods?
- If pea plants with all white seed coats are crossed, what are the possible phenotypes of their progeny?
- Pea plants with restricted yellow pods are crossed to plants that are true-breeding for inflated green pods. The F_1 are then crossed. Derive the phenotypic and genotypic ratios for the F_2 generation.
- In 1883, Alexander Graham Bell reported on records of deaf individuals in the United States, and hypothesized that advances such as sign language and schools for the deaf would lead to increased incidence of the condition resulting from an increasing number of marriages between deaf individuals. More recently, researchers reported that the number of marriages between two deaf individuals in which all children will be deaf has increased five-fold over the past century, and that the incidence of hereditary deafness has also increased. What is the most likely mode of inheritance that applies to families in which all children are born deaf?
- The MacDonalds raise Labrador retrievers. In one litter, two of eight puppies have a condition called exercise-induced collapse. After about 15 minutes of intense exercise, the dogs wobble about, develop a fever, and their hind legs collapse. The parents are healthy. What is the mode of inheritance?
- A genealogical study of 160,811 couples born from 1800 through 1965 in Iceland, an isolated population, revealed that couples who were first cousins had just as many children as couples who were second or third cousins. Is this an expected

or unexpected finding? What further information is necessary to interpret these observations?

- Draw a pedigree to depict the following family: One couple has a son and a daughter with normal skin pigmentation. Another couple has one son and two daughters with normal skin pigmentation. The daughter from the first couple has three children with the son of the second couple. Their son and one daughter have albinism; their other daughter has normal skin pigmentation.
- Chands syndrome (MIM 214350) is autosomal recessive and causes very curly hair, underdeveloped nails, and abnormally shaped eyelids. In the following pedigree, which individuals must be carriers?

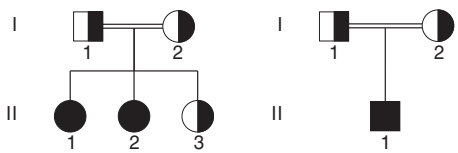


Chands syndrome

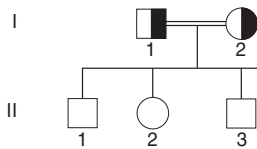
- Lorenzo has a double row of eyelashes (MIM 126300), which he inherited from his mother as a dominant trait. His maternal

grandfather is the only other relative to have it. Fatima, who has normal eyelashes, marries Lorenzo. Their first child, Nicola, has normal eyelashes. Now Fatima is pregnant again and hopes for a child with double eyelashes. What chance does the child have of inheriting double eyelashes? Draw a pedigree of this family.

11. Peeling skin syndrome (MIM 270300) causes the outer skin layer to fall off on the upper surfaces of the hands and feet. The pedigrees depict three families with this condition. What do they share that might explain the appearance of this otherwise rare condition?



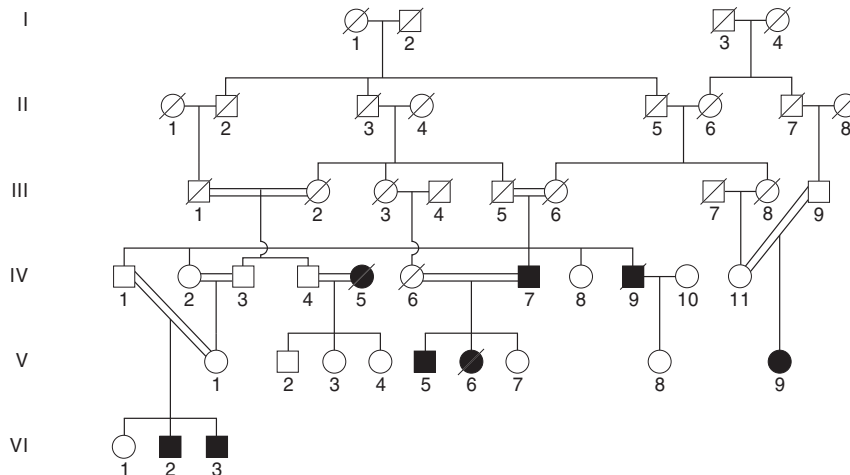
a. b.



c.

Peeling skin syndrome

12. Sclerosteosis (MIM 269500) causes overgrowth of the skull and jaws that produces a characteristic face, gigantism, facial paralysis, and hearing loss. The overgrowth of skull bones can cause severe headaches and even sudden death. In this pedigree for a family with sclerosteosis:
- What is the relationship between the individuals who are connected by slanted double lines?
 - Which individuals in the pedigree must be carriers?



Sclerosteosis

13. The child referenced in figure 4.14 who has red urine after eating beets, colored eyelids, and short fingers, is of genotype $bbHhEe$. The genes for these traits are on different chromosomes. If he has children with a woman who is a trihybrid for each of these genes, what are the expected genotypic and phenotypic ratios for their offspring?

Web Activities

- Go to the website for the National Organization for Rare Disorders. Identify an autosomal recessive disorder and an autosomal dominant disorder. Create a family for each one, and describe transmission of the disease over three generations.
- Go to the website for Gene Gateway—Exploring Genes and Genetic Disorders. Select two disorders or traits that would demonstrate independent assortment if present in the same family, and two that would not.
- Many software programs are available that construct pedigrees, even for pets. A free one is at www.hhs.gov/familyhistory. It only follows a few diseases, but is very helpful for displaying relationships. PowerPoint can also be used to construct a pedigree. Use these or other resources, or old-fashioned pencil and paper, to construct a pedigree for your own family.

Case Studies and Research Results

- On the daytime drama “The Young and the Restless,” several individuals suffer from SORAS, which stands for “soap opera rapid aging syndrome.” It is not listed in MIM. In SORAS, a young child is sent off to boarding school and returns three months later an angry teenager. In the Newman family, siblings Nicholas and Victoria aged from ages 6 and 8 years, respectively, to 16 and 18 years within a few months. Their parents, Victor and Nikki, are curiously not affected; in fact, they never seem to age at all. What is the mode of inheritance of the rapid aging disorder affecting Nicholas and Victoria, and how do you know this?
- More than a dozen recessive illnesses that are very rare in most of the world are fairly common among the Bedouin people who live in the Negev desert area of Israel. More than 65 percent of Bedouins marry their first or second cousins. This practice helped enable the group to survive a nomadic existence in the harsh environment in the past. Recently, two physicians and a geneticist have set up a service for the Bedouins that enables people wishing to marry to take genetic tests to learn if they are carriers for the same diseases. Prenatal testing has also been introduced to provide the option of terminating pregnancies that would otherwise lead to the births of children who would die of a recessive disorder in early childhood. Discuss the pros and cons of introducing genetic testing in this community, including your opinion on whether medical science should interfere with a society’s long-held cultural practices.
- The Cleaver family awaited the birth of Claudette’s puppies with great anticipation. But shortly after the standard poodle gave birth to six pups, two of them, a male and a female, developed seizures and died. One of the surviving pups, Sylvester, was bred when he was a year old to Minuette, who was an offspring of Sylvester’s father, Otis. Alas, one of Minuette’s four pups also died of the canine seizure disorder.
 - Draw a pedigree for the poodle family.
 - What is the mode of inheritance of the disorder?
 - What advice would you give to the Cleavers about successfully breeding their poodles in the future?
- A man named Grady Stiles performed in circuses as “Lobster Boy,” a condition known technically as ectrodactyly, or split-hand/split-foot syndrome (MIM 225300). Lobster Boy had only two digits on each hand and foot, but was otherwise healthy, living long enough to see the trait appear in two of his four children, adding them to his traveling act. A granddaughter has the condition too. Grady, who died in 1992, traced the family trait back six generations to a William Stiles, born in 1805. Ectrodactyly may be inherited as an autosomal dominant or an autosomal recessive trait. Which one is the more likely explanation for the trait in this family? Cite a reason for your answer.

Forensics Focus

- A woman, desperate to fill in the leaves on her family tree for an upcoming family reunion, cornered a stranger in a fast-food restaurant. Her genealogical research had identified him as a distant cousin, and she needed his DNA. He refused to cooperate, looked scared, and ran off. The woman took his discarded coffee cup and collected DNA from the spit within, which she sent on a swab to a DNA ancestry testing company. (This is a true story.)
 - Do you think the woman was justified in her action? Why or why not?
 - What are the strengths and limitations of using genealogical information (family records, word-of-mouth) versus DNA testing to construct a pedigree?
- A young woman walking to her car in a parking lot late at night was attacked and brutally raped. She kept her wits about her and was able to recall striking details of her rapist’s attire—a green silk shirt, a belt with a metal buckle that left a bruise on her abdomen, jeans, and running shoes. She also remembered that he had white skin, a shaved head, and startling blue eyes. She recalled footsteps approaching and then someone yanking the man off her, but her head was let go and hit the pavement, knocking her unconscious.

A few weeks after the rape, a naked male body washed up in a nearby river, and police found a belt buckle farther downstream that matched the pattern of the woman’s bruise. The body, however, was headless and the neck ragged, as if cut.

How can police determine the eye color of the corpse to help in identifying or ruling him out as the rapist?



Blackened nails are one sign of alkaptonuria, an inborn error of metabolism. Diseases with many symptoms may not appear to follow Mendel's laws.

CHAPTER

5

Beyond Mendel's Laws

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The Many Faces of "Black Urine Disease"

Pat Wright became aware of her alkaptonuria (MIM 203500) at age 15, when she suffered back spasms. As she got older, her spine continued to degenerate, and the cartilage in her left knee broke down. She had five children and taught for 26 years, but retired on disability at age 57.

Wright's medical records noted a "harmless" metabolic disorder. Her parents had taken her to the doctor after noticing dark-stained diapers. The pediatrician, suspecting alkaptonuria, sent a blackened diaper to geneticists, who diagnosed the disorder—but it was never explained well to the parents, who were carriers.

Wright didn't hear a name for her condition until 1997, when the surgeon who was replacing her knee found blackened cartilage around the joint. This explained the pains, stained diapers, and even the dark blue-gray color of Wright's ears, he told her, as well as her hearing loss, gallstones, and heart valve damage. The disorder disrupts an enzyme that normally breaks down the amino acid tyrosine.

A mutation that is associated with several symptoms may be confusing. In alkaptonuria, a single mutation causes several symptoms. The inability to break down tyrosine causes buildup of an acid that reacts to produce a black pigment that is deposited in urine, nails, skin, and cartilage. When urine is exposed to oxygen, it turns black.

5.1 A New View of Mendelian Genetics

Gregor Mendel's analysis of inheritance, in which "either-or" Punnett squares based on simple probabilities represent proportions of peas that are green or yellow, wrinkled or smooth, was brilliant for its time, and correct. Yet, that single-gene view is becoming greatly altered through today's genome-wide lens. Unlike Mendel's tallying of what he could see in breeding experiments, the sequencing of the human genome and tracking of how our DNA sequences vary have provided peeks at the tangled complexity of inheritance.

Single genes rarely completely control a phenotype in the way that Mendel's experiments suggested. Genes interact with each other, and with environmental influences, in intricate and myriad ways that we are just beginning to understand. That is, when transmission patterns of a visible trait do not fit autosomal recessive or autosomal dominant modes of inheritance, Mendel's laws are still operating, and the underlying genotypic ratios persist, but other factors help to mold phenotypes. Chapters 7, 9, and 11 explore some of these "outside-the-gene" influences, which include:

- other protein-encoding genes.
- other DNA sequences, such as SNPs in non-protein-encoding regions of the genome.
- RNA sequences that turn groups of genes on and off.
- epigenetic alterations of DNA, such as chemical groups that either activate or shield genes.
- environmental stimuli.

This chapter considers three general phenomena that seem to be exceptions to Mendel's laws, but are really not: gene expression, mitochondrial inheritance, and linkage.

5.2 When Gene Expression Appears to Alter Mendelian Ratios

Mendel's crosses yielded offspring that were easily distinguished: a pea is either yellow or green; a plant tall or short. For some characteristics, though, offspring classes are not in the proportions that Punnett squares or probabilities predict. In other cases, transmission patterns of a visible trait are not consistent with autosomal recessive or autosomal dominant inheritance. In these instances, either the nature of the phenotype or influences from other genes or the environment alter phenotypic ratios—that is, what is actually seen. Following are several circumstances in which phenotypic ratios appear to contradict Mendel's laws—but the laws still apply.

Lethal Allele Combinations

A genotype (allele combination) that causes death is, by definition, lethal. Death from genetic disease can occur at any stage of development or life. Tay-Sachs disease, discussed in section 2.2, for example, is lethal by age 3 or 4; Huntington

disease may not be lethal until late middle age (see figure 4.1). In a population and evolutionary sense, a lethal genotype has a more specific meaning—it causes death before the individual can reproduce, which prevents passage of genes to the next generation.

In organisms used in experiments, such as fruit flies, pea plants, or mice, lethal allele combinations remove an expected progeny class following a specific cross. For example, in a cross of heterozygous flies, homozygous recessive progeny die as embryos, leaving only heterozygous and homozygous dominant adult fly offspring. In humans, early-acting lethal alleles cause spontaneous abortion. When both a man and a woman carry a recessive lethal allele for the same gene, each pregnancy has a 25 percent chance of spontaneously aborting—this is the homozygous recessive class.

A double dose of a dominant allele may be lethal, as is the case for Mexican hairless dogs (**figure 5.1**). Inheriting one dominant allele confers the coveted hairlessness trait, but inheriting two dominant alleles is lethal to the unlucky embryo. Breeders cross hairless to hairy ("powderpuff") dogs, rather than hairless to hairless, to avoid losing the lethal homozygous dominant class—a quarter of the pups. The dogs also have missing or abnormally shaped teeth. A mutation in a gene called *forkhead box I3* (*FOXI3*) causes the condition, which is called canine ectodermal dysplasia. Humans also have the *FOXI3* gene (MIM 612351), but so far it has not been associated with a phenotype.

An example of a lethal genotype in humans is achondroplastic dwarfism (MIM 100800), which has a very distinct phenotype of a long trunk, very short limbs, and a large head with a flat face. It is inherited as an autosomal dominant trait, but is most often the result of a spontaneous (new) mutation. If two people with achondroplasia have children, each has a one in four chance of inheriting both mutant alleles. However, homozygotes are not seen, and this genotype is therefore presumed to be lethal. Observations of other species that have this mutation suggest that the homozygote would be unable to breathe because the lungs do not have room to inflate. The mutation is in the gene that encodes a receptor for a growth factor.

Multiple Alleles

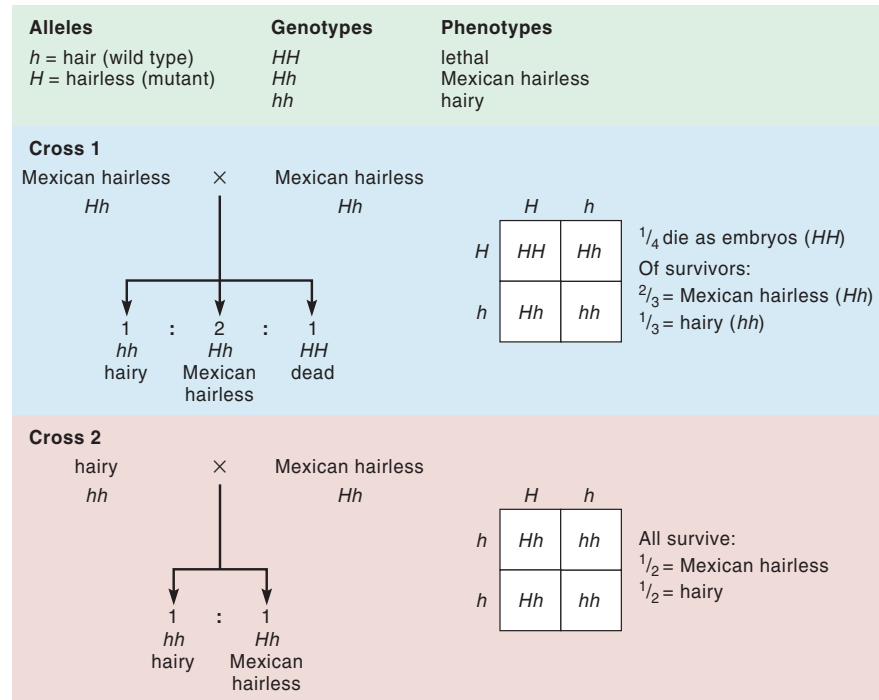
An individual has two alleles for any autosomal gene—one allele on each homolog. However, a gene can exist in more than two allelic forms in a population because it can mutate in many ways. That is, the sequence of hundreds of DNA bases that makes up a gene can be altered in many ways. Different allele combinations can produce variations in the phenotype. The more alleles, the more variations of the phenotype are possible.

It would be very useful if testing for a particular genotype could always enable physicians to predict the course of an illness. However, this is often difficult because other genes and environmental effects can modify a phenotype, a point we return to soon. Two disorders for which allele identification *can* predict severity and types of symptoms are phenylketonuria (PKU) and cystic fibrosis.

Figure 5.1 Lethal alleles. (a) This Mexican hairless dog has inherited a dominant allele that makes it hairless. Inheriting two such dominant alleles is lethal to embryos. (b) Breeders cross Mexican hairless dogs to hairy ("powderpuff") dogs to avoid dead embryos and stillbirths that represent the HH genotypic class.



a.



b.

In PKU, too little or lack of an enzyme causes the amino acid phenylalanine to build up in brain cells. Hundreds of mutant alleles pair to cause four basic phenotypes:

- classic PKU with profound mental retardation
- moderate PKU
- mild PKU
- asymptomatic PKU, with excretion of excess phenylalanine in urine

Eating a special diet extremely low in phenylalanine allows normal brain development. Knowing the allele combination can guide how strict the diet need be, and how long it must continue—generally for years.

Multiple alleles are considered in carrier testing for CF, which is done routinely in early pregnancy (see Reading 4.2). When the CF gene was discovered in 1989, researchers identified one mutant allele, $\Delta F508$, that causes about 70 percent of cases in many populations. As the allele list grew, researchers discovered that not all allele combinations cause the same symptoms. People homozygous for $\Delta F508$ have frequent, severe respiratory infections, very congested lungs, and poor weight gain. Another genotype increases susceptibility to bronchitis and pneumonia, and another causes only absence of the ductus deferens. Genetic tests probe panels of CF mutations that are the most common in a patient's ethnic group, maximizing the likelihood of detecting carriers and avoiding the cost of testing for 1,500+ alleles. If a pregnant woman has a disease-causing allele, then the father-to-be is tested, and if he has a mutant allele too, then the fetus may be tested to see if it has inherited the disease.

Different Dominance Relationships

In complete dominance, one allele is expressed, while the other isn't. In **incomplete dominance**, the heterozygous phenotype is intermediate between that of either homozygote.

In a sense, enzyme deficiencies in which a threshold level is necessary for health illustrate both complete and incomplete dominance—depending upon how one evaluates the phenotype. For example, on a whole-body level, Tay-Sachs disease displays complete dominance because the heterozygote (carrier) is as healthy as a homozygous dominant individual. However, if phenotype is based on enzyme level, then the heterozygote is intermediate between the homozygous dominant (full enzyme level) and homozygous recessive (no enzyme). Half the normal amount of enzyme is sufficient for health, which is why at the whole-person level, the wild type allele is completely dominant.

For many genes, researchers can measure the expression levels associated with various genotypes, demonstrating that even heterozygotes whose phenotypes are the same as those of homozygotes are distinctive at the biochemical level. Often, they produce half the normal amount of a protein, but this is sufficient for health.

Familial hypercholesterolemia (FH) is an example of incomplete dominance in humans that can be observed on both the molecular and whole-body levels. A person with two disease-causing alleles lacks receptors on liver cells that take up the low density lipoprotein (LDL) form of cholesterol from the bloodstream. A person with one disease-causing allele has half the normal number of receptors. Someone with two wild type (the most common) alleles has the normal number of receptors. **Figure 5.2**

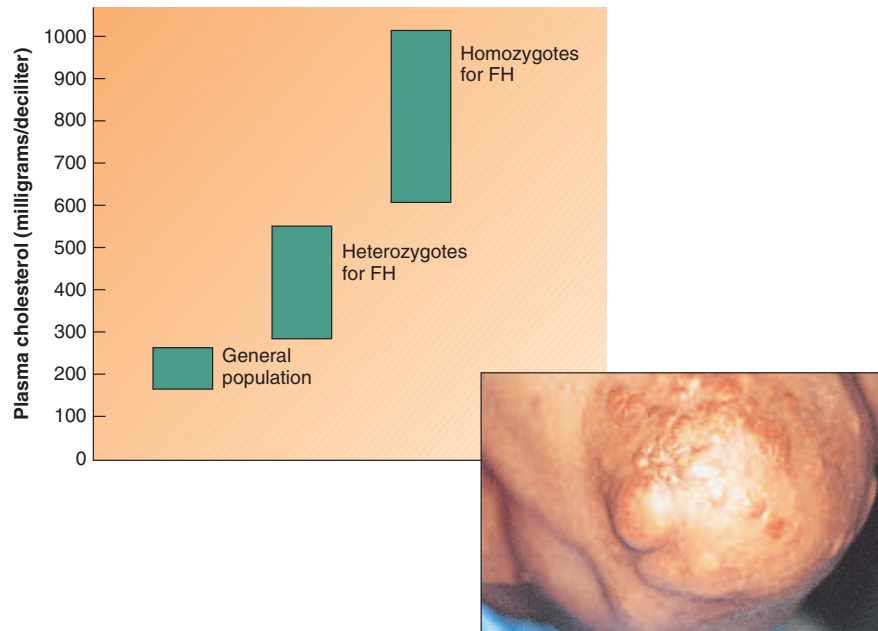


Figure 5.2 Incomplete dominance. A heterozygote for familial hypercholesterolemia (FH) has approximately half the normal number of cell surface receptors in the liver for LDL cholesterol. An individual with two mutant alleles has the severe form of FH, with liver cells that totally lack the receptors. As a result, serum cholesterol level is very high. The photograph shows cholesterol deposits on the elbow of an affected young man. Cholesterol is also deposited in joints and many other body parts.

shows how measurement of plasma cholesterol reflects these three genotypes. The phenotypes parallel the number of receptors—those with two mutant alleles die as children of heart attacks, those with one mutant allele may suffer heart attacks in young adulthood, and those with two wild type alleles do not develop this inherited form of heart disease.

Different alleles that are both expressed in a heterozygote are **codominant**. The ABO blood group is based on the expression of codominant alleles.

Blood types are determined by the patterns of molecules on the surfaces of red blood cells. Most of these molecules are proteins embedded in the plasma membrane with attached sugars that extend from the cell surface. The sugar is the antigen, which is the molecule that the immune system recognizes. People who belong to blood group A have an allele that encodes an enzyme that adds a final piece to a certain sugar to produce antigen A. In people with blood type B, the allele and its encoded enzyme are slightly different, which causes a different piece to attach to the sugar, producing antigen B. People in blood group AB have both antigen types. Blood group O reflects yet a third allele of this gene. It is missing just one DNA nucleotide, but this drastically changes the encoded enzyme in a way that robs the sugar chain of its final piece (figure 5.3). Type O red blood cells lack both A and B antigens.

The A and B alleles are codominant, and both are completely dominant to O. Considering the genotypes reveals how these interactions occur. In the past, ABO blood types have been described as variants of a gene called “*I*,” although MIM

now abbreviates the designations. The older *I* system is easier to understand. (“*I*” stands for isoagglutinin.) The three alleles are I^A , I^B , and i . People with blood type A have antigen A on the surfaces of their red blood cells, and may be of genotype $I^A I^A$ or $I^A i$. People with blood type B have antigen B on their red blood

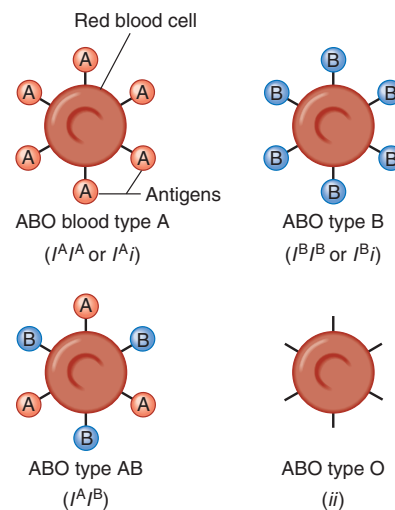


Figure 5.3 ABO blood types illustrate codominance. ABO blood types are based on antigens on red blood cell surfaces. This depiction greatly exaggerates the size of the A and B antigens. Genotypes are in parentheses.

cell surfaces, and may be of genotype $I^B I^B$ or $I^B i$. People with the rare blood type AB have both antigens A and B on their cell surfaces, and are genotype $I^A I^B$. People with blood type O have neither antigen, and are genotype ii .

Fiction plots often misuse ABO blood type terminology, assuming that a child's ABO type must match that of one parent. This is not true, because a person with type A or B blood can be heterozygous. A person who is genotype $I^A i$ and a person who is $I^B i$ can jointly produce offspring of any ABO genotype or phenotype, as **figure 5.4** illustrates.

	Type A			Type A	
	I^A	I^A		I^A	i
Type B	I^B	$I^A I^B$	AB	$I^A I^B$	AB
	i	$I^A i$	A	$I^A i$	A
	Type A			Type A	
	I^A	I^A		I^A	i
Type B	I^B	$I^A I^B$	AB	$I^A I^B$	AB
	i	$I^A i$	A	$I^A i$	A

Figure 5.4 Codominance. The I^A and I^B alleles of the I gene are codominant, but they follow Mendel's law of segregation. These Punnett squares follow the genotypes that could result when a person with type A blood has children with a person with type B blood.

Epistasis

Mendel's laws can appear not to operate when one gene masks or otherwise affects the phenotype of another. This phenomenon is called **epistasis**. It refers to interaction between different genes, not between the alleles of the same gene. A gene that affects expression of another is called a modifier gene.

In epistasis, the blocked gene is expressed normally, but the product of the modifier gene inactivates it, removes a structure needed for it to contribute to the phenotype, or otherwise counteracts its effects. An obvious example of epistasis is the hairless gene in dogs. Genes that color dog hairs have no effect if there are no hairs. An epistatic interaction seen in many species is albinism, in which one gene blocks the action of genes whose products confer color.

A more complex example of epistasis is a blood type called the Bombay phenotype. It results from an interaction between a gene called H and the I gene that confers ABO blood type. The H gene controls the placement of a molecule to which antigens A and B attach on red blood cell surfaces. In a person of genotype hh , that molecule isn't made, so the A and B antigens have no way to attach to the red blood cell surface. The A and B antigens fall off—and the person tests as type O blood, although any ABO genotype is possible.

Epistasis can explain why siblings who inherit the same disorder can suffer to differing degrees. One study examined siblings who both inherited spinal muscular atrophy 1

(MIM 253300), in which nerves cannot signal muscles. The muscles weaken and atrophy, usually proving fatal in early childhood. The mutation encodes an abnormal protein that shortens axons, which are the extensions on nerve cells that send messages. Some siblings who inherited the SMA mutation never developed symptoms. They can thank another gene, *plastin 3* (MIM 300131), which increases production of the cytoskeletal protein actin that extends axons. Because these children inherited the ability to make extra long axons, the axon-shortening effects of SMA are not harmful.

Penetrance and Expressivity

The same allele combination can produce different degrees of a phenotype in different individuals because a gene does not act alone. That is, every gene that is expressed functions against a backdrop of the expression of other genes, as well as effects of environmental influences such as nutrition, exposure to toxins, and stress. This is why two individuals who have the most severe CF genotype may have different clinical experiences. One person may be much sicker because she also inherited genes predisposing her to develop asthma and respiratory allergies. Even identical twins with the same genetic disease may be affected to different degrees due to environmental influences.

Many single-gene traits and illnesses have distinctive phenotypes, despite all of these influences. The terms *penetrance* and *expressivity* describe degrees of expression of a single gene. **Penetrance** refers to the all-or-none expression of a genotype; **expressivity** refers to severity or extent.

An allele combination that produces a phenotype in everyone who inherits it is completely penetrant—this is very rare. Huntington disease (see *Bioethics: Choices for the Future* in chapter 4) is nearly completely penetrant—almost all people who inherit the mutant allele will develop symptoms if they live long enough.

A genotype is incompletely penetrant if some individuals do not express the phenotype (have no symptoms). Polydactyly (see figure 1.5) is incompletely penetrant. Some people who inherit the dominant allele have more than five digits on a hand or foot. Yet others who we know have the allele because they have an affected parent and child have ten fingers and ten toes. Penetrance is described numerically. If 80 of 100 people who inherit the dominant polydactyly allele have extra digits, the genotype is 80 percent penetrant.

A phenotype is variably expressive if symptoms vary in intensity among different people. One person with polydactyly might have an extra digit on both hands and a foot, but another might have just one extra fingertip. Polydactyly is both incompletely penetrant and variably expressive.

It is hard to imagine how other genes or the environment can influence the numbers of fingers or toes. For familial hypercholesterolemia, variable expressivity reflects greater influence of other genes that regulate lipid levels in the blood and the environment (see figure 5.2). FH heterozygotes develop heart disease due to high serum cholesterol in middle adulthood. Healthful diet and exercise habits as well as lipid-lowering drugs can delay symptom onset.

Pleiotropy

A single-gene disorder with many symptoms, or a gene that controls several functions or has more than one effect, is termed **pleiotropic**. Such conditions can be difficult to trace through families because people with different subsets of symptoms may appear to have different disorders. A classic case of pleiotropy is porphyria variegata, an autosomal dominant disease that affected several members of the royal families of Europe (**figure 5.5**).

King George III ruled England during the American Revolution. At age 50, he first experienced abdominal pain and constipation, followed by weak limbs, fever, a fast pulse, hoarseness, and dark red urine. Next, nervous system signs and symptoms began, including insomnia, headaches, visual problems, restlessness, delirium, convulsions, and stupor. His confused and racing thoughts, combined with actions such as ripping off his wig and running about naked while at the peak of a fever, convinced court observers that the king was mad. Just as Parliament was debating his ability to rule, he recovered.

But the king's ordeal was far from over. He relapsed 13 years later, then again 3 years after that. Always the symptoms appeared in the same order, beginning with abdominal pain, fever, and weakness, and progressing to nervous system symptoms. Finally, an attack in 1811 placed George in a prolonged stupor, and the Prince of Wales dethroned him. George III lived for several more years, experiencing further episodes.

In George III's time, doctors were permitted to do very little to the royal body, and their diagnoses were based on what the king told them. Twentieth-century researchers found that porphyria variegata caused George's red urine. It is one of several types of porphyrias, which result from deficiency of any of several enzymes required to manufacture heme. The king's disorder arises from lack of enzyme #7 in the heme pathway shown in **figure 5.6**. Heme is part of hemoglobin, the molecule that carries oxygen in the blood and imparts the red color. In the disease, a part of heme called a porphyrin ring is routed

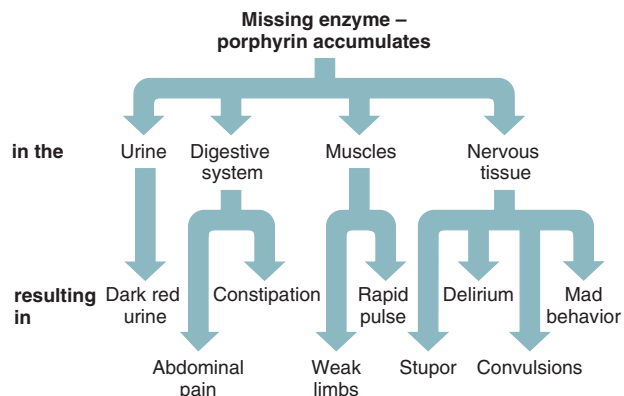


Figure 5.5 Pleiotropy. King George III suffered from the autosomal dominant disorder porphyria variegata—and so did several other family members. Because of pleiotropy, the family's varied illnesses and quirks appeared to be different, unrelated disorders. Symptoms appear every few years in a particular order.

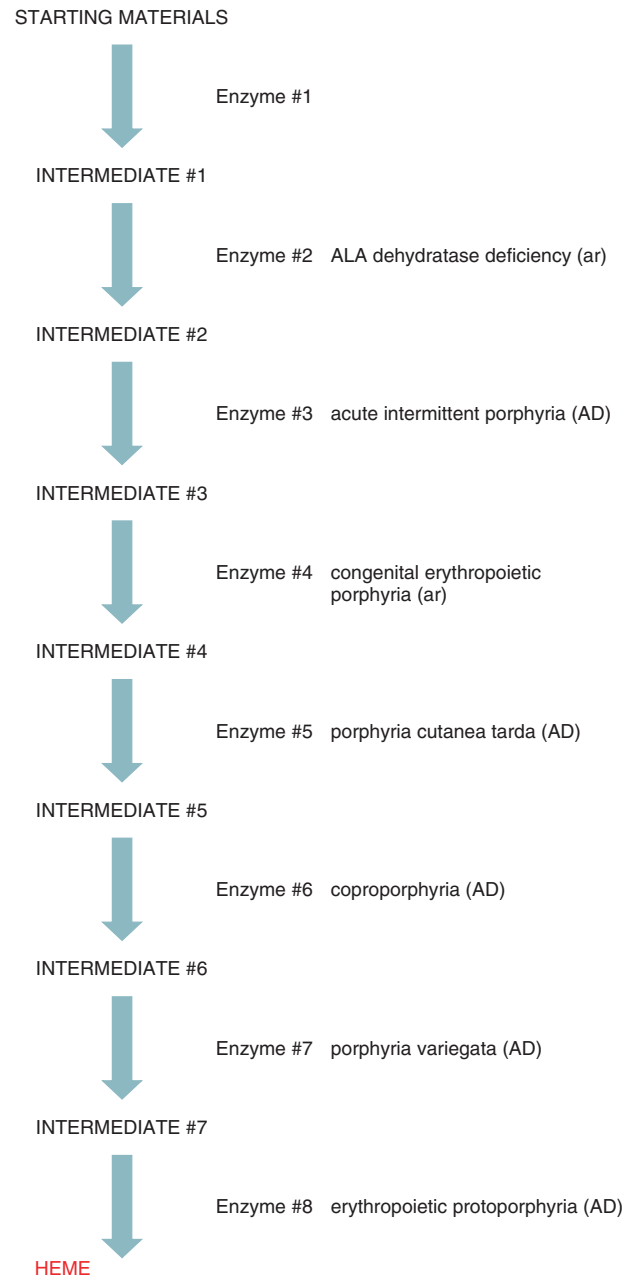


Figure 5.6 The porphyrias. Errors in the heme biosynthetic pathway cause seven related, yet distinct, diseases. In each disorder, the intermediate biochemical that a deficient enzyme would normally affect builds up. The excess is excreted in the urine or accumulates in blood, feces, or inside red blood cells, causing symptoms. People with various porphyria-related symptoms may have inspired the vampire and werewolf legends, including reddish teeth, pink urine, excess hair, and photosensitivity (avoidance of daylight).

into the urine instead of being broken down and metabolized in cells. Porphyrin also builds up and attacks the nervous system.

Examination of physicians' reports on George's royal relatives showed that the disorder was thought to be several different illnesses. Today, porphyria variegata remains rare, and is often misdiagnosed as a seizure disorder. Unfortunately, some seizure medications and anesthetics worsen symptoms. Ironically, treatment may have worsened King George III's disease. Medical records and hair analysis indicate that a medicine based on the element antimony was forced upon the king in the madhouse. Antimony was often contaminated with arsenic, and arsenic inactivates several of the enzymes in the heme biosynthetic pathway!

On a molecular level, pleiotropy occurs when a single protein affects different body parts, participates in more than one biochemical reaction, or has different effects in different amounts. Consider Marfan syndrome. The most common form of this autosomal dominant condition is a defect in an elastic connective tissue protein called fibrillin (MIM 134797). The protein is abundant in the lens of the eye, in the aorta (the largest artery in the body, leading from the heart), and in the bones of the limbs, fingers, and ribs. The symptoms are lens dislocation, long limbs, spindly fingers, and a caved-in chest. The most serious symptom is a life-threatening weakening in the aorta, which can suddenly burst. If the weakening is detected early, a synthetic graft can replace the section of artery wall.

Certain proteins that form a structure in the eye called lens crystallins beautifully illustrate pleiotropy. If in low abundance as single molecules, these proteins are metabolic enzymes, functioning in many cell types. At higher abundance, however, they join and form crystallins, which aggregate to create a transparent lens whose physical properties enable it to focus incoming light on the retina. These multifunctional lens crystalline proteins are well-studied in a variety of species, including insects, jellyfish, and vertebrates.

Genetic Heterogeneity

When different genes produce the same phenotype, a phenomenon called **genetic heterogeneity**, it may appear that Mendel's laws are not operating. For example, 132 forms of hearing loss are transmitted as autosomal recessive traits. If a man who is homozygous for a hearing loss gene on one chromosome has a child with a woman who is homozygous for a hearing loss gene on a different chromosome, then the child would not be deaf, because he or she would be heterozygous for both hearing-related genes. The different forms of hearing loss reflect the many ways that this sense can be genetically impaired.

Discovering additional genes that can cause a known disorder is happening more as the human genome is analyzed, and can have practical repercussions. Consider osteogenesis imperfecta, in which abnormal collagen causes children's bones to break easily (see figure 3.21). Often when a child is brought to the hospital and past fractures are discovered, child abuse is suspected. A test for the most common type of osteogenesis imperfecta (MIM 166210) can rule this out. However, some parents

who insisted there was no abuse did not have the common mutation. They were cleared of charges in 2006 when researchers discovered a rare mutation in a different gene (MIM 610854). This gene normally encodes an enzyme that adds a small chemical group to collagen, enabling it to function.

Genetic heterogeneity can occur when genes encode enzymes that catalyze the same biochemical pathway, or different proteins that are part of the pathway. **Reading 5.1** describes a common genetically heterogeneous condition—Alzheimer disease.

Phenocopies

An environmentally caused trait that appears to be inherited is a **phenocopy**. Such a trait can either produce symptoms that resemble those of a known single-gene disorder or mimic inheritance patterns by affecting certain relatives. For example, the limb birth defect caused by the drug thalidomide, discussed in chapter 3, is a phenocopy of the inherited illness phocomelia. Physicians recognized the environmental disaster when they began seeing many children born with what looked like the very rare phocomelia. A birth defect caused by exposure to a teratogen was more likely than a sudden increase in incidence of a rare inherited disease.

An infection can be a phenocopy. Children who have AIDS may have parents who also have the disease, but these children acquired AIDS by viral infection, not by inheriting a gene. A phenocopy caused by a highly contagious infection can seem to be inherited if it affects more than one family member.

Sometimes, common symptoms may resemble those of an inherited condition until medical tests rule heredity out. For example, an underweight child who has frequent colds may show some signs of cystic fibrosis, but may instead suffer from malnutrition. Negative test results for several common CF alleles would alert a physician to look for another cause.

The Human Genome Sequence Adds Perspective

As researchers continue to identify and describe the genes sequenced in the human genome project, it is becoming clear that phenomena once considered to complicate single-gene inheritance aren't rare, and they may be common. As a result, terms such as *epistasis* and *genetic heterogeneity* are beginning to overlap and blur. Consider Marfan syndrome. Most affected individuals have a mutation in the fibrillin gene. However, some people with the syndrome instead have a mutation in the gene that encodes the transforming growth factor beta receptor (TGF β R) (MIM 190181). Fibrillin and TGF β R are part of the same biochemical pathway. The conditions fit the definition of genetic heterogeneity because mutations in different genes cause identical symptoms. Yet they are also epistatic because a mutation in TGF β R blocks the activity of fibrillin.

Gene interactions also underlie penetrance and expressivity, once thought to be strictly a characteristic of a particular gene. Even genes that do not directly interact, in space or time, can affect each other's expression. This is the case for

Reading 5.1

The Genetic Roots of Alzheimer Disease

"What is that thing for, that you put in your ear?" asked 72-year-old Ginny for the fifth time in half an hour.

"Mom, don't you know you've asked me that several times? It gets cell phone calls," answered her son, trying not to become annoyed.

"No, I've never asked you that before." She paused, looking puzzled.

"What did you say it is?"

In the following months, Ginny's short-term memory declined further. She could rarely concentrate enough to finish reading a newspaper article, or follow a long conversation. In the grocery store she had trouble finding items she had been buying for decades. Aware of her growing deficits, she became depressed. Finally, her son suggested she have a complete neurological exam. By the time she could see a physician, other signs had emerged. Ginny couldn't recall her zip code or the name of the small town where she grew up. Sometimes she couldn't remember where things belong—she once put a cantaloupe in the bathtub, and still had trouble with that contraption her son put in his ear to receive phone calls.

The doctor sent Ginny for a brain scan, which supported her diagnosis of "mild cognitive impairment." It could be the beginnings of Alzheimer disease. She started Ginny on a cholinesterase inhibitor drug to slow breakdown of the neurotransmitter acetylcholine, which sends messages in the brain. She also prescribed an anti-depressant, which revived Ginny enough so that she was more willing to leave

her apartment. Despite these treatments, over the next several years Ginny would continue to lose cognitive function—thinking, reasoning, learning, and communicating—and would one day no longer recognize her son or her friends. Eventually, she would lose the ability to speak and smile, and would cease walking, not because her legs wouldn't function, but because she would forget how. Towards the end, she would not even be able to sit up or lift her head. Yet, even in her final days, in fleeting moments when the haze would seem to lift from her eyes, the old Ginny would return.

In Alzheimer disease, certain brain parts—the amygdala (seat of emotion) and hippocampus (the memory center)—become mired in two types of protein. Amyloid precursor protein is cut into abnormally sized amyloid beta peptides that aggregate to form "plaques" outside brain cells. Tau proteins (MIM 157140) become encased in too much phosphate, which makes them adhere into "tangles" inside brain cells, blocking their normal function of binding to microtubules (**figure 1**). The telltale plaques and tangles, present to a lesser extent in everyone, could cause the symptoms or, more likely, result from them.

Fewer than 1 percent of Alzheimer's cases are familial (inherited), caused by mutations in any of at least three genes (**table 1**). A variant of a fourth gene, *APOE4*, increases the risk of developing a late-onset form three-fold in a heterozygote and 15-fold in a homozygote. Other genes raise the risk, too. A variant of a gene called *CALHM1* raises risk to about a tenth of the extent of *APOE4*, and genome-wide association studies have identified five places in the genome that may harbor other risk genes. The sporadic (non-inherited) cases begin later in life and reflect the cumulative effects of inheriting several common susceptibility alleles. Alzheimer disease is, clearly, genetically heterogeneous.

Table 1 Genes Associated with Alzheimer Disease

	MIM	Chromosome	Mechanism
Causative Gene			
Amyloid precursor protein (<i>APP</i>)	104760	21	Abnormally sized pieces aggregate outside brain cells.
Presenilin 1	607822 104311	14	Forms part of secretase (enzyme) that cuts APP.
Presenilin 2	606889 600759	1	Forms part of secretase (enzyme) that cuts APP.
Risk Gene			
Apolipoprotein E4 (<i>APOE4</i>)	104310 107741	19	Apolipoprotein is cut into abnormally sized pieces that add phosphates to tau protein, making it accumulate and impairing microtubule binding.

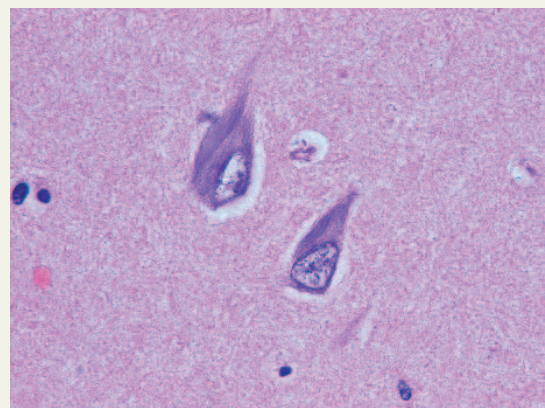


Figure 1 The dark dots in these two brain neurons are "neurofibrillary tangles" of tau protein, characteristic of Alzheimer disease.

Huntington disease, described in chapter 4, in which cells in a certain part of the brain die, typically beginning in young adulthood. Siblings who inherit the exact same HD mutation may differ in the number of cells that they have in the affected brain area, thanks to variants of other genes that affected the division rate of neural stem cells in the brain during embryonic development. As a result, an individual who inherits HD, but also extra brain cells, might develop symptoms much later in life than a brother or sister who does not have such a built-in reserve supply. If the delay is long enough that death comes from another cause, HD would then be nonpenetrant.

DNA microarrays that reveal gene expression patterns in different tissues are painting detailed portraits of pleiotropy, showing that inherited disorders may affect more tissues or organs than are obvious as symptoms. Finally, more cases of genetic heterogeneity are being discovered as researchers identify genes with redundant or overlapping functions.

Table 5.1 summarizes phenomena that appear to alter single-gene inheritance. Our definitions and designations are changing as improving technology enables us to describe and differentiate disorders in greater detail. Phenomena such as variable expressivity, incomplete penetrance, epistasis, pleiotropy, and genetic heterogeneity, once considered unusual characteristics of single genes, are turning out to be the norm.

Gregor Mendel derived the two laws of inheritance working with traits conferred by genes located on different chromosomes in the nucleus. When genes do not conform to these conditions, however, the associated traits may not appear in Mendelian ratios. The remainder of this chapter considers two types of gene transmission that do not fulfill the requirements for single-gene inheritance.

Key Concepts

1. A lethal genotype does not appear as a progeny class.
2. In incomplete dominance, the heterozygote phenotype is intermediate between those of the homozygotes; in codominance, two different alleles for the same gene are expressed.
3. In epistasis, one gene influences expression of another.
4. Genotypes vary in penetrance and expressivity of the phenotype.
5. A gene with more than one expression is pleiotropic.
6. In genetic heterogeneity, different genes cause the same phenotype.
7. A trait caused by the environment but resembling a known genetic trait or occurring in certain family members is a phenocopy.

5.3 Mitochondrial Genes

The basis of the law of segregation, that both parents contribute genes equally to offspring, does not apply for genes in mitochondria (see figure 2.7). The thousands of mitochondria in each human cell contain several copies of a “mini-chromosome” that carries just 37 genes. mtDNA-encoded genes act in the mitochondrion, but the organelle also requires the activities of certain genes from the nucleus.

The inheritance patterns and mutation rates for mitochondrial genes differ from those for genes in the nucleus.

Table 5.1 Factors That Alter Single-Gene Phenotypic Ratios

Phenomenon	Effect on Phenotype	Example
Lethal alleles	A phenotypic class does not survive to reproduce.	Spontaneous abortion
Multiple alleles	Many variants or degrees of a phenotype occur.	Cystic fibrosis
Incomplete dominance	A heterozygote's phenotype is intermediate between those of two homozygotes.	Familial hypercholesterolemia
Codominance	A heterozygote's phenotype is distinct from and not intermediate between those of the two homozygotes.	ABO blood types
Epistasis	One gene masks or otherwise affects another's phenotype.	Bombay phenotype
Penetrance	Some individuals with a particular genotype do not have the associated phenotype.	Polydactyly
Expressivity	A genotype is associated with a phenotype of varying intensity.	Polydactyly
Pleiotropy	The phenotype includes many symptoms, with different subsets in different individuals.	Porphyria variegata
Phenocopy	An environmentally caused condition has symptoms and a recurrence pattern similar to those of a known inherited trait.	Infection
Genetic heterogeneity	Different genotypes are associated with the same phenotype.	Hearing impairment

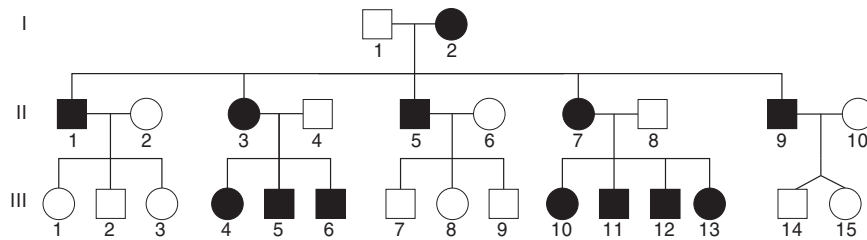


Figure 5.7 Inheritance of mitochondrial genes. Mothers pass mitochondrial genes to all offspring. Fathers do not transmit mitochondrial genes because sperm only very rarely contribute mitochondria to fertilized ova. If mitochondria from a male do enter, they are destroyed.

Mitochondrial genes are maternally inherited. They are passed only from an individual's mother because sperm almost never contribute mitochondria when they fertilize an oocyte. In the rare instances when mitochondria from sperm enter an oocyte, they are usually selectively destroyed early in development. Pedigrees that follow mitochondrial genes therefore show a woman passing the trait to all her children, while a male cannot pass the trait to any of his (figure 5.7).

DNA in the mitochondria differs functionally from DNA in the nucleus in several ways (table 5.2 and figure 5.8). Mitochondrial DNA does not cross over. It mutates faster than DNA in the nucleus for two reasons: it has fewer ways to repair DNA (discussed in chapter 12), and the mitochondrion is the site of energy reactions that produce oxygen free radicals that damage DNA. Also unlike nuclear DNA, mtDNA is not wrapped in proteins, nor are genes "interrupted" by DNA sequences that do not encode protein. Finally, a cell has one nucleus but many mitochondria—and each mitochondrion harbors several copies of its chromosome. Mitochondria with different alleles for the same gene can reside in the same cell.

Mitochondrial Disorders

Mitochondrial genes encode proteins that participate in protein synthesis and energy production. Twenty-four of the 37 genes encode RNA molecules (22 transfer RNAs and 2 ribosomal RNAs) that help assemble proteins. The other 13 mitochondrial genes encode proteins that function in cellular respiration,

which is the process that uses energy from digested nutrients to synthesize ATP, the biological energy molecule.

Several diseases result from mutations in mitochondrial genes. They are called mitochondrial myopathies and have specific names, but news reports often lump them together as "mitochondrial disease." Symptoms arise from tissues whose cells have many mitochondria, such as skeletal muscle. It isn't surprising that major symptoms are great fatigue, weak and flaccid muscles, and intolerance to exercise. Skeletal muscle fibers appear "red and ragged" when stained and viewed under a light microscope, their abundant abnormal mitochondria visible beneath the plasma membrane.

A defect in an energy-related gene can produce symptoms other than fatigue. This is the case for Leber optic atrophy (MIM 535000), which impairs vision. First described in 1871 and its maternal transmission noted, this disorder was associated in 1988 with a mitochondrial mutation that impairs cellular energy reactions. Symptoms usually begin in early adulthood with a loss of central vision. Eyesight worsens and color vision vanishes as the central portion of the optic nerve degenerates. Chapter 20 describes gene therapy for this condition.

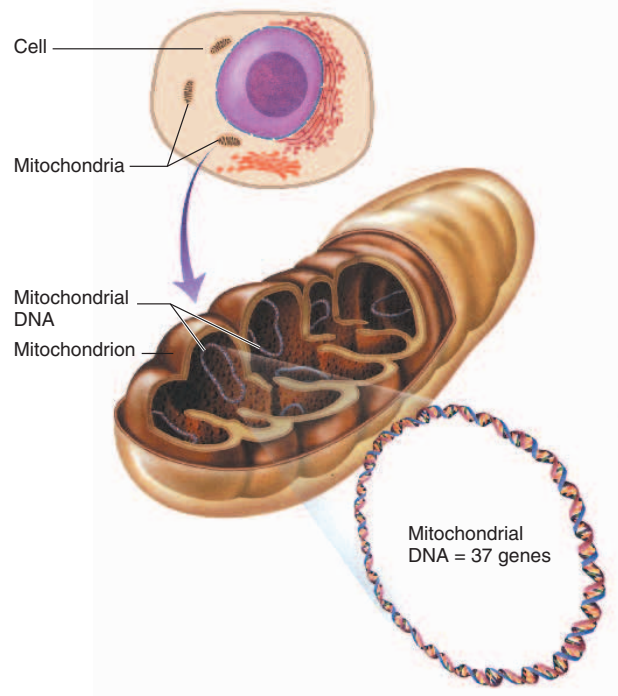


Figure 5.8 Mitochondrial DNA. A mitochondrion contains several rings of DNA. Different alleles can reside on different copies of the mitochondrial chromosome. A cell typically has thousands of mitochondria, each of which has many copies of its "mini-chromosome."

Table 5.2	Features of Mitochondrial DNA
	No crossing over
	Fewer types of DNA repair
	Inherited from the mother only
	Many copies per mitochondrion and per cell
	High exposure to oxygen free radicals
	No histones (DNA-associated proteins)
	Genes not interrupted

A mutation in a mitochondrial gene that encodes a tRNA or rRNA can be devastating because it impairs the organelle's ability to manufacture proteins. Consider what happened to Lindzy S., a once active and articulate dental hygienist. In her forties, Lindzy gradually began to slow down at work. She heard a buzzing in her ears and developed difficulty talking and walking. Then her memory began to fade in and out, she became lost easily in familiar places, and her conversation made no sense. Her condition worsened, and she developed diabetes, seizures, and pneumonia and became deaf and demented. She was finally diagnosed with MELAS, which stands for "mitochondrial myopathy encephalopathy lactic acidosis syndrome" (MIM 540000). Lindzy died. Her son and daughter will likely develop the condition because they inherited her mitochondria.

A woman with a mitochondrial disorder can avoid transmitting it if mitochondria from a healthy woman's oocyte are injected into her oocytes. Such a bolstered oocyte is fertilized in a laboratory dish by the partner's sperm, and the zygote is implanted in her uterus. Several dozen children, apparently free of mitochondrial disease, have been born from this technique.

About 1 in 200 people has a mutation in a mitochondrial gene that could cause disease. However, mitochondrial diseases are very rare, apparently because of a weeding-out process during egg formation. Such a mutation may disrupt energy acquisition, which dooms the oocyte harboring it.

Heteroplasmy

The fact that a cell contains many mitochondria makes possible a rare condition called **heteroplasmy**. In this state a particular mutation is in some mitochondrial chromosomes, but not others. At each cell division, the mitochondria are distributed at random into daughter cells. Over time, the chromosomes within a mitochondrion tend to be all wild type or all mutant for any particular gene. But different mitochondria can have different

alleles predominating. As an oocyte matures, the number of mitochondria drops from about 100,000 to 100 or fewer. If the woman is heteroplasmic for a particular mutation, by chance, she can produce an oocyte that has mostly mitochondria that are wild type for that gene, mostly mitochondria that have the mutation, or anything in between (**figure 5.9**). In this way, a woman who does not have a mitochondrial disorder, because the mitochondria bearing the mutation are either rare or not abundant in affected cell types, can nevertheless pass the associated condition to a child.

Heteroplasmy has several consequences for the inheritance of mitochondrial phenotypes. Expressivity may vary widely among siblings, depending upon how many mutation-bearing mitochondria were in the oocyte that became each brother or sister. Severity of symptoms reflects which tissues have cells whose mitochondria bear the mutation. This is the case for a family with Leigh syndrome (MIM 256000), which affects the enzyme that directly produces ATP. Two boys died of the severe form of the disorder because the brain regions that control movement rapidly degenerated. Another sibling was blind and had central nervous system degeneration. Several relatives, however, suffered only mild impairment of their peripheral vision. The more severely affected family members had more brain cells that received the mutation-bearing mitochondria.

The most severe mitochondrial illnesses are heteroplasmic. This is presumably because *homoplasmy*—when all mitochondria bear the mutant allele—too severely impairs protein synthesis or energy production for embryonic development to complete. Often, severe heteroplasmic mitochondrial disorders do not produce symptoms until adulthood because it takes many cell divisions, and therefore years, for a cell to receive enough mitochondria bearing mutant alleles to cause symptoms. This is why Leber optic atrophy usually does not affect vision until adulthood.

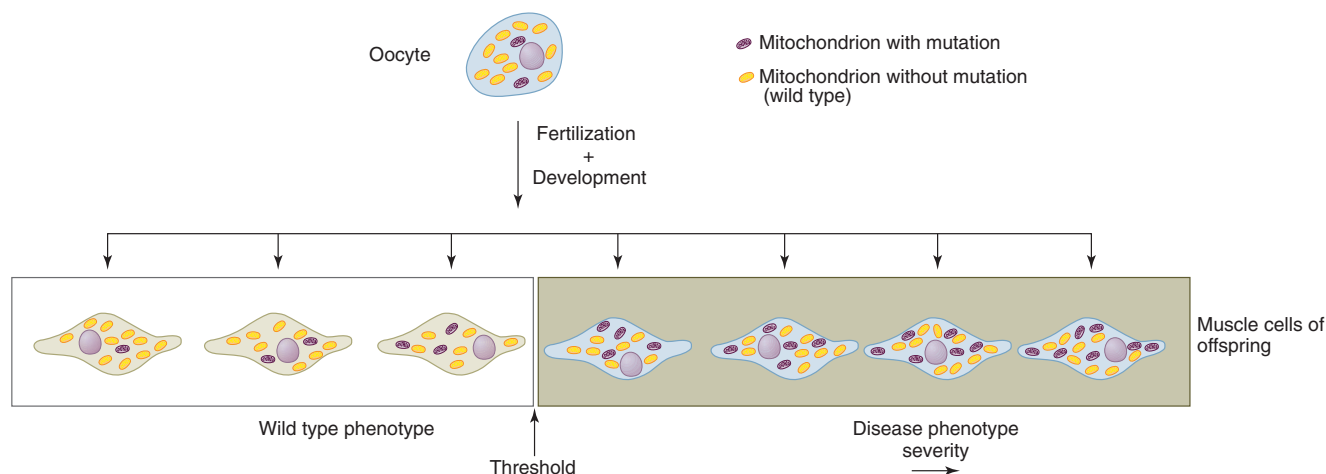


Figure 5.9 Mitochondrial inheritance. Mitochondria and their genes are only passed from the mother. Cells have many mitochondria. If an oocyte is heteroplasmic, differing numbers of copies of a mitochondrial mutation may be transmitted. The phenotype reflects the proportion of mitochondria bearing the mutation.

Mitochondrial DNA Reveals the Past

Interest in mtDNA extends beyond the medical. mtDNA provides a powerful forensic tool used to link suspects to crimes, identify war dead, and support or challenge historical records. mtDNA is used in forensics because it is more likely to remain after extensive damage, because cells have many copies of it.

Sequencing mtDNA identified the son of Marie Antoinette and Louis XVI, who supposedly died in prison at age 10. In 1845, the boy was given a royal burial, but some people thought the buried child was an imposter. His heart had been stolen at the autopsy, and through a series of bizarre events, wound up, dried out, in the possession of the royal family. Recently, researchers compared mtDNA sequences from cells in the boy's heart to corresponding sequences in heart and hair cells from Marie Antoinette (her decapitated body identified by her fancy underwear), two of her sisters, and living relatives Queen Anne of Romania and her brother. The mtDNA evidence showed that the buried boy was indeed the prince, Louis XVII.

Key Concepts

1. Mitochondrial genes are maternally inherited and mutate rapidly. A cell contains many mitochondria, which have many copies of the mitochondrial genome.
2. Mitochondrial genes encode RNAs or proteins that function in protein synthesis or energy metabolism.
3. In heteroplasmy, cells contain mitochondria that have different alleles of a gene.

5.4 Linkage

Most of the traits that Mendel studied in pea plants were conferred by genes on different chromosomes. When genes are close to each other on the same chromosome, they usually do not segregate at random during meiosis and therefore do not support Mendel's predictions. Instead, genes close on a chromosome are packaged into the same gametes and are said to be "linked" (**figure 5.10**). Linkage has this very precise meaning in genetics. The term is popularly used in a much more general sense to mean any association between two events or observations.

Linkage refers to the transmission of genes on the same chromosome. Linked genes do *not* assort independently and do *not* produce Mendelian ratios for crosses tracking two or more genes. Understanding and using linkage as a mapping tool has been critical in identifying disease-causing genes, and helped pave the way for genome-wide association studies.

Discovery in Pea Plants

William Bateson and R. C. Punnett first observed the unexpected ratios indicating linkage in the early 1900s, again in pea plants. They crossed true-breeding plants with purple flowers

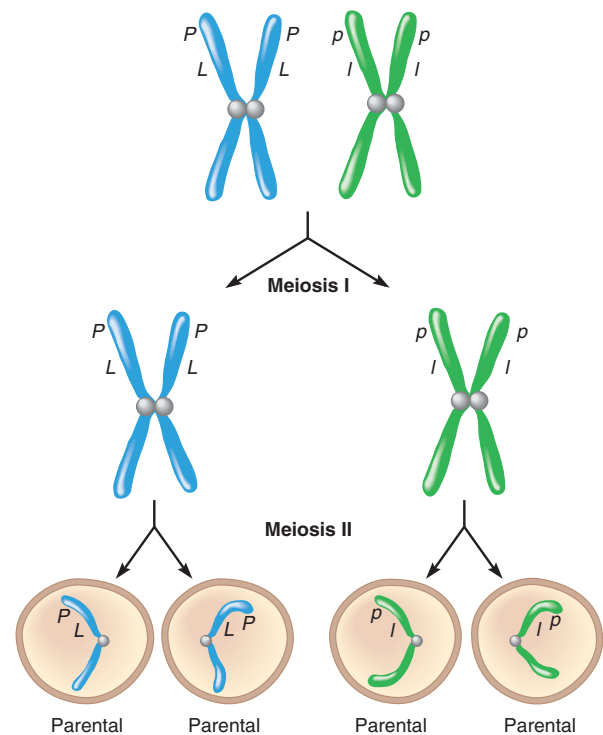
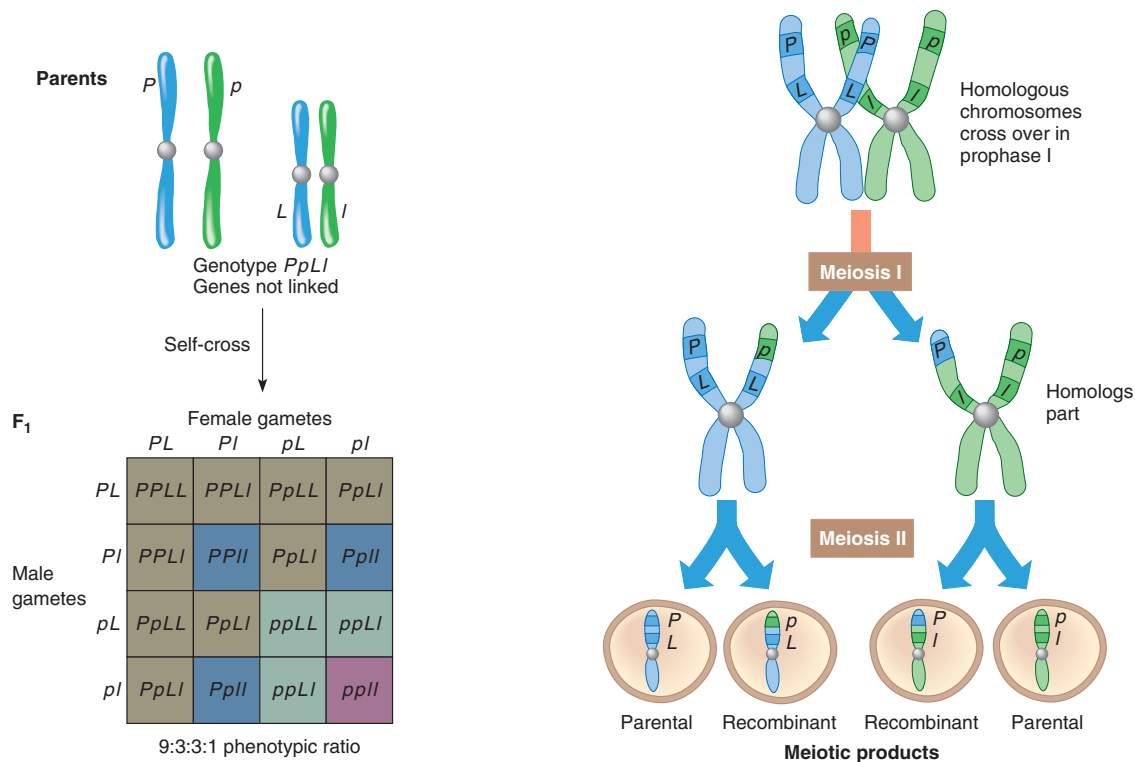


Figure 5.10 Inheritance of linked genes. Genes linked closely to one another are usually inherited together when the chromosome is packaged into a gamete.

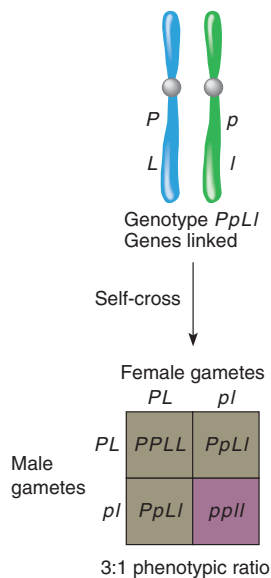
and long pollen grains (genotype *PPLL*) to true-breeding plants with red flowers and round pollen grains (genotype *ppll*). The plants in the next generation, of genotype *PpLl*, were then self-crossed. But this dihybrid cross did not yield the expected 9:3:3:1 phenotypic ratio that Mendel's second law predicts (**figure 5.11**).

Bateson and Punnett noticed that two types of third-generation peas, those with the parental phenotypes *P_L_* and *ppll*, were more abundant than predicted, while the other two progeny classes, *ppL_* and *P_ll*, were less common (the blank indicates that the allele can be dominant or recessive). The more prevalent parental allele combinations, Bateson and Punnett hypothesized, could reflect genes that are transmitted on the same chromosome and that therefore do not separate during meiosis. The two less common offspring classes could also be explained by a meiotic event—crossing over. Recall that this is an exchange between homologs that mixes up maternal and paternal gene combinations without disturbing the sequence of genes on the chromosome (**figure 5.12**).

Progeny that exhibit this mixing of maternal and paternal alleles on a single chromosome are called **recombinant**. *Parental* and *recombinant* are relative terms. Had the parents in Bateson and Punnett's crosses been of genotypes *ppL_* and *P_ll*, then *P_L_* and *ppll* would be recombinant rather than parental classes.



a.



b.

Figure 5.11 Expected results of a dihybrid cross. (a)

Unlinked genes assort independently. The gametes represent all possible allele combinations. The expected phenotypic ratio of a dihybrid cross is 9:3:3:1. (b) If genes are linked, only two allele combinations are expected in the gametes. The phenotypic ratio is 3:1, the same as for a monohybrid cross.

Figure 5.12 Crossing over disrupts linkage. The linkage between two genes may be interrupted if the chromosome they are on crosses over with its homolog between the two genes. Crossing over packages recombinant groupings of the genes into gametes.

Two other terms describe the configurations of linked genes in dihybrids. Consider a pea plant with genotype $PpLl$. These alleles can be part of the chromosomes in either of two ways. If the two dominant alleles are on one chromosome and the two recessive alleles on the other, the genes are in “*cis*.” In the opposite configuration, with one dominant and one recessive allele on each chromosome, the genes are in “*trans*” (figure 5.13). Whether alleles in a dihybrid are in *cis* or *trans* is important in distinguishing recombinant from parental progeny classes in specific crosses.

Linkage Maps

As Bateson and Punnett were discovering linkage in peas, geneticist Thomas Hunt Morgan and his coworkers at Columbia University were doing the same using the fruit fly *Drosophila melanogaster*. They assigned genes to relative positions on chromosomes and compared progeny class sizes to assess whether traits were linked. They soon saw that the pairs of traits fell into four groups. Within each group, crossed dihybrids did not produce offspring classes according to Mendel's second law. Also, the number of linkage groups—four—matched the number of chromosome pairs in the fly. Coincidence? No. The traits fell into four groups because their genes are inherited together on the same chromosome.

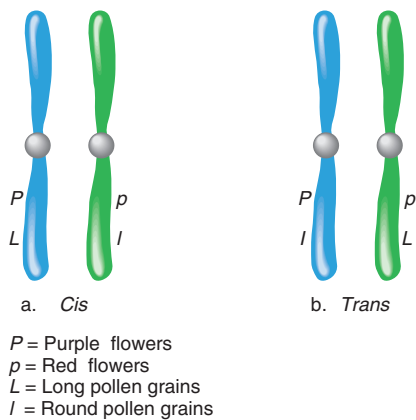


Figure 5.13 Allele configuration is important. Parental chromosomes can be distinguished from recombinant chromosomes only if the allele configuration of the two genes is known—they are either in *cis* (a) or in *trans* (b).

The genius of the work on linkage in fruit flies was that the researchers translated their data into actual maps depicting positions of genes on chromosomes. Morgan wondered why the sizes of the recombinant classes varied for different genes. In 1911, Morgan's undergraduate student, Alfred Sturtevant, proposed that the farther apart two genes are on a chromosome, the more likely they are to cross over simply because more physical distance separates them (figure 5.14).

The correlation between crossover frequency and the distance between genes is used to construct **linkage maps**. These diagrams show the order of genes on chromosomes and the relative distances between them. The distance is represented using “map units” called centimorgans, where 1 centimorgan equals 1 percent recombination. These units are used today to construct SNP maps. They provide an estimate of genetic distance along a chromosome.

The frequency of a crossover between any two linked genes is inferred from the proportion of offspring that are recombinant. Frequency of recombination is based on the percentage of meiotic

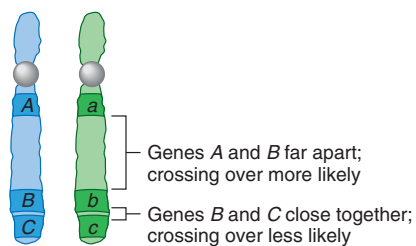


Figure 5.14 Breaking linkage. Crossing over is more likely to occur between the widely spaced linked genes A and B, or between A and C, than between the more closely spaced linked genes B and C, because there is more room for an exchange to occur.

divisions that break the linkage between—that is, separate—two parental alleles. Genes at opposite ends of the same chromosome cross over often, generating a large recombinant class. Genes lying very close on the chromosome would only rarely be separated by a crossover. The probability that genes on opposite ends of a chromosome cross over approaches the probability that, if on different chromosomes, they would independently assort—about 50 percent. **Figure 5.15** illustrates this distinction.

The situation with linked genes (or SNPs) can be compared to a street lined with stores on both sides. There are more places to cross the street between stores at opposite ends on opposite sides than between two stores in the middle of the block on opposite sides of the street. Similarly, more crossovers, or progeny with recombinant genotypes, are seen when two genes are farther apart on the same chromosome.

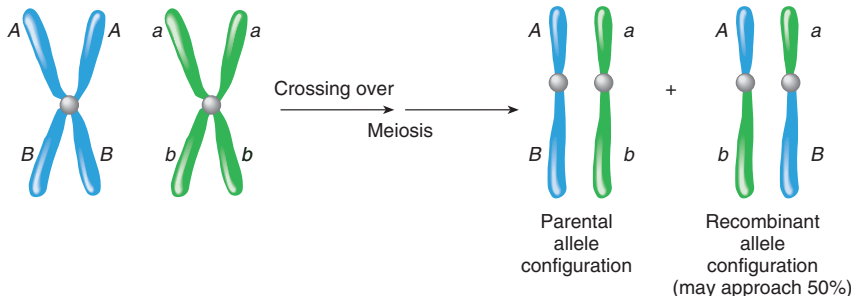
As the twentieth century progressed, geneticists in Columbia University's “fly room” mapped several genes on all four chromosomes, as researchers in other labs assigned many genes to the human X chromosome. Localizing genes on the X chromosome was easier than doing so on the autosomes, because X-linked traits follow an inheritance pattern that is distinct from the one followed by all autosomal genes. In human males, with their single X chromosome, recessive alleles on the X are expressed. We return to this point in chapter 6.

By 1950, geneticists began to think about mapping genes on the 22 human autosomes. To start, a gene must be matched to its chromosome. This became possible when people with a particular inherited condition or trait also had a specific chromosome abnormality. Matching phenotypes to chromosomal variants, a field called cytogenetics, is the subject of chapter 13.

In 1968, researchers assigned the first human gene to an autosome. R. P. Donohue was observing chromosomes in his own white blood cells when he noticed a dark area consistently located near the centromere of one member of his largest chromosome pair (chromosome 1). He examined chromosomes from several family members for the dark area, noting also whether each family member had a blood type called Duffy. (Recall that blood types refer to the patterns of sugars on red blood cell surfaces.) Donohue found that the Duffy blood type was linked to the chromosome variant. He could predict a relative's Duffy blood type by whether or not the chromosome had the telltale dark area.

Finding a chromosomal variation and using it to detect linkage to another gene was a valuable but rare achievement. More often, researchers used the experiments like those Sturtevant did on flies—calculating percent recombination (crossovers) between two genes with known locations on a chromosome. However, because humans do not have hundreds of offspring, nor do they produce a new generation every 10 days, obtaining sufficient data to establish linkage relationships requires observing the same traits in many families and pooling the information. Today, even though we know the human genome sequence, linkage remains a powerful tool to track disease-associated genes.

Linked Genes



Nonlinked Genes

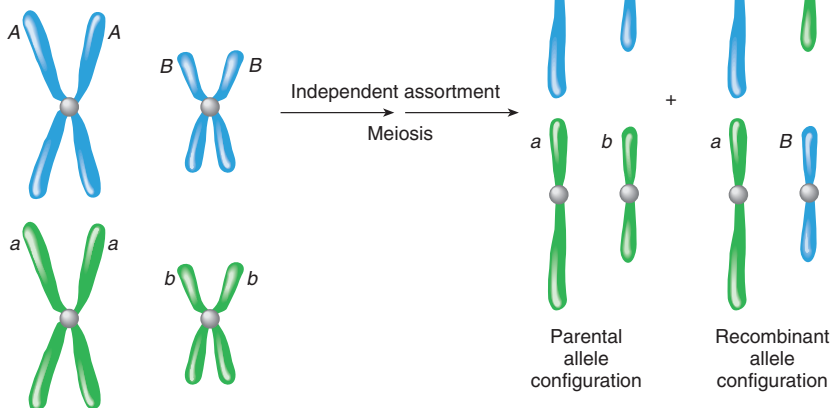


Figure 5.15 Linkage versus nonlinkage (independent assortment). When two genes are widely separated on a chromosome, the likelihood of a crossover is so great that the recombinant class may approach 50 percent—which may appear to be the result of independent assortment.

⊕ Solving a Problem

Linkage

As an example of determining the degree of linkage by percent recombination, consider the traits of Rh blood type and a form of anemia called elliptocytosis (MIM 111700) (MIM 130500). An Rh⁺ phenotype corresponds to genotypes *RR* or *Rr*. (This is simplified.) The anemia corresponds to genotypes *EE* or *Ee*.

In 100 one-child families, one parent is Rh negative with no anemia (*rre*), and the other parent is Rh positive with anemia (*RrEe*), and the *R* and *E* (or *r* and *e*) alleles are in *cis*. Of the 100 offspring, 96 have parental genotypes (*re/re* or *RE/re*) and four are recombinants for these two genes (*Re/re* or *rE/re*). Percent recombination is therefore 4 percent, and the two linked genes are 4 centimorgans apart.

Consider another pair of linked genes in humans. Nail-patella syndrome (MIM 161200) is a rare autosomal dominant trait that causes absent or underdeveloped fingernails and toenails, and painful arthritis in the knee and elbow joints. The gene is 10 map units from the *I* gene that determines

the ABO blood type, on chromosome 9. Geneticists determined the map distance by pooling information from many families. The information can be used to predict genotypes and phenotypes in offspring, as in the following example.

Greg and Susan each have nail-patella syndrome. Greg has type A blood. Susan has type B blood. What is the chance that their child inherits normal nails and knees and type O blood? A genetic counselor deduces their allele configurations using information about the couple's parents (figure 5.16).









Greg's mother has nail-patella syndrome and type A blood. His father has normal nails and type O blood. Therefore, Greg must have inherited the dominant nail-patella syndrome allele (*N*) and the *I*^A allele from his mother, on the same chromosome. We know this because Greg has type A blood and his father has type O blood—therefore, he couldn't have gotten the *I*^A allele from his father. Greg's other chromosome 9 must carry the alleles *n* and *i*. His alleles are therefore in *cis*.

Susan's mother has nail-patella syndrome and type O blood, and so Susan inherited *N* and *i* on the same chromosome. Because her father has normal nails and type B blood, her homolog from him bears alleles *n* and *I*^B. Her alleles are in *trans*.

Determining the probability that Susan and Greg's child could have normal nails and knees and type O blood is the easiest question the couple could ask. The only way this genotype can arise from theirs is if an *ni* sperm (which occurs with a frequency of 45 percent, based on pooled data) fertilizes an *ni* oocyte (which occurs 5 percent of the time). The result—according to the product rule—is a 2.25 percent chance of producing a child with the *nnii* genotype.

Calculating other genotypes for their offspring is more complicated, because more combinations of sperm and oocytes could account for them. For example, a child with nail-patella syndrome and type AB blood could arise from all combinations that include *I*^A and *I*^B as well as at least one *N* allele (assuming that *NN* has the same phenotype as *Nn*).

The Rh blood type and elliptocytosis, and nail-patella syndrome and ABO blood type, are examples of linked gene pairs. A linkage map begins to emerge when percent recombination is known between all possible pairs of three or more linked genes, just as a road map with more landmarks provides more information on distance and direction. Consider genes *x*,

	Greg		Susan	
Phenotype	nail-patella syndrome, type A blood		nail-patella syndrome, type B blood	
Genotype	$NnI^A i$		$NnI^B _$	
Allele configuration	$\frac{N}{n} \frac{I^A}{i}$		$\frac{N}{n} \frac{i}{I^B}$	
Gametes:	sperm	frequency	oocytes	
Parental		45%		
		45%		
Recombinants		5%		
		5%		

N = nail-patella syndrome
 n = normal

Figure 5.16 Inheritance of nail-patella syndrome. Greg inherited the N and I^A alleles from his mother; that is why the alleles are on the same chromosome. His n and i alleles must therefore be on the homolog. Susan inherited alleles N and i from her mother, and n and I^B from her father. Population-based probabilities are used to calculate the likelihood of phenotypes in the offspring of this couple. Note that in this figure, map distances are known and are used to predict outcomes.

y , and z (**figure 5.17**). If the percent recombination between x and y is 10, between x and z is 4, and between z and y is 6, then the order of the genes on the chromosome is x - z - y . This is the only order of the three genes that accounts for the percent recombination data.

Knowing the percent recombination between linked genes was useful in crudely ordering them on genetic maps. Understanding the structure of DNA, and then sequencing the human genome, revealed an unexpected complexity in linkage mapping: crossing over is not equally likely across the genome. Some DNA sequences are nearly always inherited together, more often than would be predicted from their frequency in the population. This nonrandom association between DNA sequences is called **linkage disequilibrium** (LD). The human genome consists of many “LD” blocks where stretches of alleles stick together, interspersed with areas where crossing over is prevalent. Chapter 7 discusses use of **haplotypes** (LD blocks) to track genes of interest in populations.

% recombination between gene pairs	
$x - y$	10
$x - z$	4
$z - y$	6

x	10	y		
----- -----		-----		
x	4	z	6	y
----- ----- -----				

Figure 5.17 Recombination mapping. If we know the percent recombination between all possible pairs of three genes, we can determine their relative positions on the chromosome.

From Linkage to Genome-Wide Associations

Gene mapping has had an interesting history. In the first half of the twentieth century, gene maps for nonhuman organisms, such as fruit flies, were constructed based on recombination frequencies between pairs of visible traits. In the 1950s, linkage data on traits in humans began to accumulate. At first, it was mostly a few visible or measurable traits linked to blood types or blood proteins.

In 1980 came a great stride in linkage mapping. Researchers began using DNA sequences near genes of interest as landmarks called **genetic markers**. These markers need not necessarily encode a protein that causes a phenotype—they might be DNA sequence differences that alter where a DNA cutting enzyme cuts, or differing numbers of short repeated sequences of DNA with no obvious function, or single nucleotide polymorphisms (SNPs) (see figure 7.10). The term “genetic marker” is used popularly to mean any DNA sequence that is associated with a particular phenotype, usually one affecting health.

The sequencing of the human genome has taken DNA mapping to a new level, but built on the old idea of linkage. Instead of relying on a few neighbors of protein-encoding genes in families, **genome-wide association studies** scan the genomes of thousands of individuals who have a particular phenotype for DNA sequences that they share, but are much rarer in people who do not have the trait (see figure 7.9). The “signposts” are SNPs and repeats. The underlying assumption is that these shared genome regions will point the way toward genes that control or contribute to the phenotype. The great value of the new types of markers is that they can shed light on the biological bases of common, multifactorial disorders. With this powerful new tool, it is becoming clear that Mendel was not wrong, but that single genes are only part of the picture of how our DNA shapes us. Genome-wide association studies are discussed in detail in chapter 7, and **table 5.3** reviews types of genetic markers mentioned throughout the book.

Computers tally how often genes and their markers are inherited together. The “tightness” of linkage between a marker and a gene of interest is represented as a LOD score, which stands for “logarithm of the odds.” A LOD score indicates the likelihood that particular crossover frequency data indicate linkage, rather than the inheritance of two alleles by chance.

A LOD score of 3 or greater signifies linkage. It means that the observed data are 1,000 (10^3) times more likely to have occurred if the two DNA sequences (a disease-causing allele and its marker) are linked than if they reside on different chromosomes and just happen to often be inherited together by chance. It is somewhat like deciding whether two coins tossed together 1,000 times always come up both heads or both tails by chance, or because they are taped together side by side in that position,

Table 5.3 Types of Genetic Markers

Genetic Element	Description	Example/Application
Chromosomal	Trait appears with unusual chromosome	Duffy blood type and dark area of chromosome 1
RFLPs	DNA cutting site	Transmission of Huntington disease in a family (Reading 22.1)
VNTRs	Repeats	Forensics (DNA profiling) (table 14.4)
STRs	Repeats	Forensics (DNA profiling) (table 14.4)
SNPs	Single-base differences	Genome-wide association studies to identify health-related genes (figure 7.9)
CNVs	Copy number variants	Cholesterol level (figure 12.11)

as linked genes are. If the coins land with the same sides up in all 1,000 trials, it indicates they are very likely attached.

Before sequencing of the human genome, genetic markers were used to predict which individuals in some families were most likely to have inherited a particular disorder, before symptoms began. Such tests are no longer necessary, because tests directly detect disease-causing genes. However, genetic markers are still used to distinguish parts of chromosomes. In pedigrees, marker designations are sometimes placed beneath the traditional

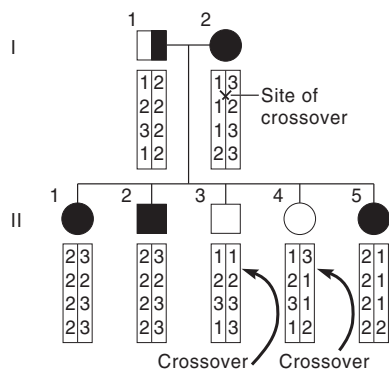


Figure 5.18 Haplotypes. The numbers in bars beneath pedigree symbols enable researchers to track specific chromosome segments with markers. Disruptions of a marker sequence indicate crossover sites.

symbols to further describe chromosomes. Such a panel of linked markers, a haplotype, is a set of DNA sequences inherited together on the same chromosome due to linkage disequilibrium.

Consider the haplotypes in a family with cystic fibrosis in **figure 5.18**. Each set of numbers beneath a symbol represents a haplotype. Knowing the haplotype of individual I-2 reveals which chromosome in parent I-1 contributes the mutant allele. Because Mr. II-2 received haplotype 3233 from his affected mother, his other haplotype, 2222, comes from his father. Since Mr. II-2 is affected and his father is not, the father must be a heterozygote, and 2222 must be the haplotype linked to the mutant CFTR allele.

Today, new associations between DNA sequences and traits are reported daily. Genetics has indeed evolved from an obscure science to a robust source of knowledge about ourselves.

Key Concepts

1. Genes on the same chromosome are linked. They are inherited in different patterns than are unlinked genes.
2. Crosses involving linked genes produce a large parental class and a small recombinant class (caused by crossing over).
3. The farther apart two genes are on a chromosome, the more likely they are to cross over. Linkage maps translate crossover frequency into relative distances between genes on a chromosome.
4. Cytogenetic abnormalities revealed the first linkage associations.
5. Linkage disequilibrium is a linkage combination that is stronger than that predicted by gene frequencies in a population. Genetic markers were used in early linkage mapping and today in genome-wide association studies.
6. Linkage maps reflect the percent recombination between linked genes. LOD scores describe the tightness of linkage. Haplotypes indicate linked DNA sequences, and reflect a variety of marker types.

Summary

5.1 A New View of Mendelian Genetics

1. Single genes do not act as independently as Mendel's experiments suggested.
2. Changes in gene expression, inheritance of mitochondrial genes, and linkage may seem to oppose Mendel's laws, but do not.

5.2 When Gene Expression Appears to Alter Mendelian Ratios

3. Homozygosity for lethal recessive alleles stops development before birth, eliminating an offspring class.

- A gene can have multiple alleles because its sequence can deviate in many ways. Different allele combinations may produce different variations of the phenotype.
- Heterozygotes of **incompletely dominant** alleles have phenotypes intermediate between those associated with the two homozygotes. **Codominant** alleles are both expressed in the phenotype.
- In **epistasis**, one gene affects the phenotype of another.
- An incompletely **penetrant** genotype is not expressed in all individuals who inherit it. Phenotypes that vary in intensity among individuals are variable in **expressivity**.
- Pleiotropic** genes have several expressions.
- In **genetic heterogeneity**, two or more genes specify the same phenotype.
- A **phenocopy** is a characteristic that appears to be inherited but is environmentally caused.

5.3 Mitochondrial Genes

- Only females transmit mitochondrial genes; males can inherit such a trait but cannot pass it on.
- Mitochondrial genes do not cross over, and they mutate more frequently than nuclear DNA.
- The 37 mitochondrial genes encode tRNA, rRNA, or proteins involved in protein synthesis or energy reactions.
- Many mitochondrial disorders are **heteroplasmic**, with mitochondria in a single cell harboring different alleles.

5.4 Linkage

- Genes on the same chromosome are **linked** and, unlike genes that independently assort, produce many individuals with parental genotypes and a few with **recombinant** genotypes.
- Linkage maps** depict linked genes. Researchers can examine a group of known linked DNA sequences (a **haplotype**) to follow the inheritance of certain chromosomes.
- Knowing whether linked alleles are in *cis* or *trans*, and using crossover frequencies from pooled data, one can predict the probabilities that certain genotypes will appear in progeny.
- Genetic linkage maps assign distances to linked genes based on crossover frequencies. Today many genetic markers spanning the genome are used to compare large groups of individuals, one of which has a particular phenotype. These **genome-wide association studies** are used in discovering genetic variants that contribute to multifactorial conditions.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

- Explain how each of the following phenomena can disrupt Mendelian phenotypic ratios.
 - lethal alleles
 - multiple alleles
 - incomplete dominance
 - codominance
 - epistasis
 - complete penetrance
 - variable expressivity
 - pleiotropy
 - a phenocopy
 - genetic heterogeneity
- How does the relationship between dominant and recessive alleles of a gene differ from epistasis?
- Why can transmission of an autosomal dominant trait with incomplete penetrance look like autosomal recessive inheritance?
- How does inheritance of ABO blood type exhibit both complete dominance and codominance?
- How could two people with albinism have a child who has normal skin pigment?
- How do the porphyrias exhibit variable expressivity, pleiotropy and genetic heterogeneity?
- How can epistasis explain incomplete penetrance?
- The lung condition emphysema may be caused by lack of an enzyme, or by smoking. Which cause is a phenocopy?
- List three ways that mtDNA differs from DNA in a cell's nucleus.
- Describe why inheritance of mitochondrial DNA and linkage are exceptions to Mendel's laws.
- How does a pedigree for a maternally inherited trait differ from one for an autosomal dominant trait?
- If researchers could study pairs of human genes as easily as they can study pairs of genes in fruit flies, how many linkage groups would they detect?
- Describe three types of genetic markers.
- The popular media often use words that have precise meanings in genetics, but more general common meanings. Explain the two types of meanings of "linked" and "marker."

Applied Questions

- For each of the diseases described in situations *a* through *i*, indicate which of the following phenomena (A–H) is at work. More than one may apply.
 - lethal alleles
 - multiple alleles
 - epistasis
 - incomplete penetrance
 - variable expressivity
 - pleiotropy
 - a phenocopy
 - genetic heterogeneity
 - A woman has severe neurofibromatosis type 1. She has brown spots on her skin and several large tumors beneath her skin. A genetic test shows that her son has inherited the disease-causing autosomal dominant allele, but he has no symptoms.
 - A man would have a widow's peak, if he wasn't bald.
 - A man and woman have six children. They also had two stillbirths—fetuses that died shortly before birth.
 - Mutations in any of at least three genes cause familial ALS.
 - A woman with dark brown skin uses a bleaching cream with a chemical called hydroquinone that darkens her fingertips and ears, just like alkaptonuria.
 - In Labrador retrievers, the *B* allele confers black coat color and the *b* allele brown coat color. The *E* gene controls the expression of the *B* gene. If a dog inherits the *E* allele, the coat is golden no matter what the *B* genotype is. A dog of genotype *ee* expresses the *B* (black) phenotype.
 - Two parents are heterozygous for genes that cause albinism, but each gene specifies a different enzyme in the biochemical pathway for skin pigment synthesis. Their children thus do not face a 25 percent risk of having albinism.
 - Alagille syndrome (MIM 118450), in its most severe form, prevents the formation of ducts in the gallbladder, causing liver damage. Affected children also usually have heart murmurs, unusual faces, a line in the eye, and butterfly-shaped vertebrae. Such children often have one otherwise healthy parent who has a heart murmur, unusual face, and butterfly vertebrae.
 - Two young children in a family have terribly decayed teeth. Their parents think it is genetic, but the true cause is a babysitter who puts them to sleep with juice bottles in their mouths.
- If many family studies for a particular autosomal recessive condition reveal fewer affected individuals than Mendel's law predicts, the explanation may be either incomplete penetrance or lethal alleles. How might you use haplotypes to determine which of these two possibilities is the cause?
- A man who has type O blood has a child with a woman who has type A blood. The woman's mother has AB blood, and her father, type O. What is the probability that the child is of blood type
 - O
 - A
 - B
 - AB?
- Enzymes are used in blood banks to remove the A and B antigens from blood types A and B. This makes the blood type O.
 - Does this alter the phenotype or the genotype?
 - Removing the A and B antigens from red blood cells is a phenocopy of what genetic phenomenon?
- Ataxia-oculomotor apraxia syndrome (MIM 208920), which impairs the ability to feel and move the limbs, usually begins in early adulthood. The molecular basis of the disease is impairment of ATP production in mitochondria, but the mutant gene is in the nucleus of the cells. Would this disorder be inherited in a Mendelian fashion? Explain your answer.
- What is the chance that Greg and Susan, the couple with nail-patella syndrome, could have a child with normal nails and type AB blood?
- A gene called secretor (MIM 182100) is located 1 map unit from the *H* gene that confers the Bombay phenotype on chromosome 19. Secretor is dominant, and a person of either genotype *SeSe* or *Sese* secretes the ABO and H blood type antigens in saliva and other body fluids. This secretion, which the person is unaware of, is the phenotype. A man has the Bombay phenotype and is not a secretor. A woman does not have the Bombay phenotype and is a secretor. She is a dihybrid whose alleles are in *cis*. What is the chance that their child will have the same genotype as the father?
- In prosopagnosia (MIM 610382), a person has "face blindness"—he or she cannot identify individuals by their faces. It is inherited as an autosomal dominant trait, and affects people to different degrees. Some individuals learn early in life—perhaps from a parent—to identify people by other features, such as voice or style of dress, and so appear not to have the condition. Only a small percentage of cases are inherited; most are the result of stroke or brain injury. www.faceblind.org offers tests to help you imagine what it is like not to be able to recognize faces—not even your own. Which of the following does face blindness demonstrate? Explain your choices.
 - incomplete penetrance
 - variable expressivity
 - pleiotropy
 - phenocopy
- Many people who have the "iron overload" disease hereditary hemochromatosis (MIM 235200; see section 20.2) are homozygous for a variant of the *C282Y* gene. How would you determine the penetrance of this condition?
- A Martian creature called a gazook has 17 chromosome pairs. On the largest chromosome are genes for three traits—round or square eyeballs (*R* or *r*); a hairy or smooth tail (*H* or *h*); and 9 or 11 toes (*T* or *t*). Round eyeballs, hairy tail, and 9 toes are dominant to square eyeballs, smooth tail, and 11 toes. A trihybrid male has

offspring with a female who has square eyeballs, a smooth tail, and 11 toes on each of her three feet. She gives birth to 100 little gazooks, who have the following phenotypes:

- 40 have round eyeballs, a hairy tail, and 9 toes
- 40 have square eyeballs, a smooth tail, and 11 toes
- 6 have round eyeballs, a hairy tail, and 11 toes
- 6 have square eyeballs, a smooth tail, and 9 toes

- 4 have round eyeballs, a smooth tail, and 11 toes
 - 4 have square eyeballs, a hairy tail, and 9 toes
- a. Draw the allele configurations of the parents.
 - b. Identify the parental and recombinant progeny classes.
 - c. What is the crossover frequency between the *R* and *T* genes?

Web Activities

11. Go to the Family Village website. Family Village is a clearinghouse for disease information. Click on library. Explore the diseases, and identify one that exhibits pleiotropy.
12. Go to the United Mitochondrial Disease Foundation website and describe the phenotype of a mitochondrial disorder.
13. Browse the National Center for Biotechnology Information (NCBI) site, and list three sets of linked genes. Consult MIM to describe the trait or disorder that each specifies.
14. Use MIM to identify a genetically heterogeneous condition, and explain why this description applies.
15. For some of the porphyrias, attacks are precipitated by an environmental trigger. Using MIM, describe factors that can trigger an attack of any of the following:
 - a. acute intermittent porphyria
 - b. porphyria cutanea tarda
 - c. coproporphyrinuria
 - d. porphyria variegata
 - e. erythropoietic protoporphyria

Case Studies and Research Results

16. Shiloh Winslow is deaf. In early childhood, she began having fainting spells, especially when she became excited. When she fainted while opening Christmas gifts, her parents took her to the hospital, where doctors said, as they had in the past, that there wasn't a problem. As the spells continued, Shiloh became able to predict the attacks, telling her parents that her head hurt beforehand. Her parents took her to a neurologist, who checked Shiloh's heart and diagnosed long QT syndrome with deafness, a severe form of inherited heartbeat irregularity (see Reading 2.2). Ten different genes can cause long QT syndrome. The doctor told them of a case from 1856: a young girl, called at school to face the headmaster for an infraction, became so agitated that she dropped dead. The parents were not surprised; they had lost two other children to great excitement.

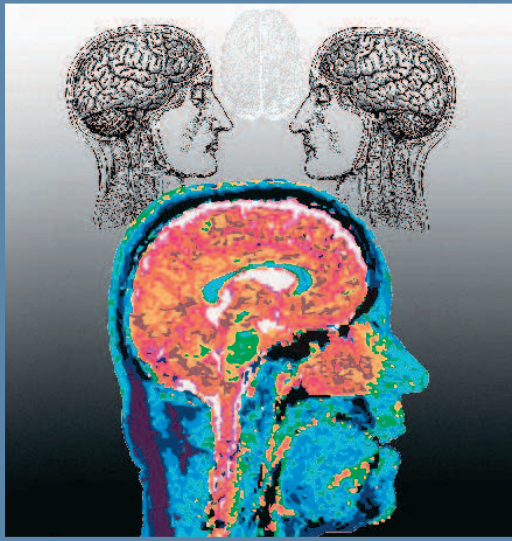
The Winslows visited a medical geneticist, who discovered that each parent had a mild heartbeat irregularity that did not produce symptoms. Shiloh's parents had normal hearing. Shiloh's younger brother Pax was also hearing-impaired and suffered night terrors, but had so far not fainted during the day. Like Shiloh, he had the full syndrome. Vivienne, still a baby, was also tested. She did not have either form of the family's illness; her heartbeat was normal.

Today, Shiloh and Pax are treated with beta blocker drugs, and each has an implantable defibrillator to correct a potentially fatal heartbeat. Shiloh's diagnosis may have saved her brother's life.

 - a. Which of the following applies to the condition in this family?
 - i. genetic heterogeneity
 - ii. pleiotropy
 - iii. variable expressivity
 - iv. incomplete dominance
 - v. a phenocopy
 - b. How is the inheritance pattern of this form of long QT syndrome similar to that of familial hypercholesterolemia?
 - c. How is it possible that Vivienne did not inherit either the serious or asymptomatic form of the illness?
 - d. Do the treatments for the condition affect the genotype or the phenotype?
17. Barnabas Collins has congenital erythropoietic porphyria, and his wife Angelique is a carrier of ALA dehydratase deficiency. What is the chance that if they have a child, he or she will have a porphyria?

Forensics Focus

18. "Earthquake McGoon" was 32 years old when the plane he was piloting over north Vietnam was hit by groundfire on May 6, 1954. Of the five others aboard, only two survived. McGoon, actually named James B. McGovern, was well known for his flying in World War II, and for his jolliness. Remains of a man about his height and age at death were discovered in late 2002, but could not be identified by dental records. However, DNA sampled from a leg bone enabled forensic scientists to identify him. Describe the type of DNA likely analyzed, and what further information was needed to make the identification.



The fact that autism and schizophrenia have opposite symptoms suggests that perhaps they can result from abnormal expression of the same gene or genes, which in turn may reflect which parent transmits the expressed allele. The imprinted brain hypothesis suggests that males can transmit autism, and females can transmit schizophrenia. It is an intriguing idea—but so far, just that.

CHAPTER

6

Matters of Sex

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A Controversial Hypothesis: Mental Illness, Mom, and Dad

Mental illnesses have been extremely difficult to classify. They run in families yet not in predictable patterns, and attempts to pin such disorders as autism, depression, bipolar disorder, and schizophrenia on specific genes or environmental influences have been mired in complexity. A sociologist from London and an evolutionary biologist from Canada have turned fresh eyes to the challenge of classifying mental disorders.

Their view, which they call the “imprinted brain hypothesis,” places the disorders on a continuum, rather than in the separate bins of the psychiatrist’s diagnostic manual. Sociologist Christopher Badcock and biologist Bernard Crespi argue that the disorders stem not so much from which gene variants patients inherit, but whether the alleles come from the father or mother—a phenomenon called genomic imprinting.

The idea did not come from DNA sequencing, but from simple observation: Dr. Badcock noticed that the way that children with autism avoid making eye contact with people and speak of themselves in the third person seems opposite the way that people with paranoid schizophrenia imagine they are being followed by others and have deluded ideas of their own importance. This view echoes a genetic phenomenon called imprinting, in which certain genes are physically blocked from being expressed, by being covered in chemical methyl (CH₃) groups, if they come from one parent but not if they come from the other. Imprinting is well-studied in certain pairs of illnesses that have symptoms similar to autism if a particular gene variant comes from the father, yet symptoms more like schizophrenia or a mood disorder if inherited from the mother.

The imprinted brain hypothesis has galvanized much new research on the underpinnings of these common and disturbing disorders. Although it is likely that the conditions are indeed caused and influenced by numerous genetic and environmental factors, researchers can now investigate whether certain genes cause autism when underexpressed, or schizophrenia when overexpressed. Perhaps most importantly, this new view of the brain may shed light on the biochemical balance that lies behind the range of moods and behaviors that we consider to be normal.

6.1 Our Sexual Selves

Whether we are male or female is enormously important in our lives, affecting our relationships, how we think and act, and how others perceive us. Gender is, at one level, dictated by genes but it is also layered with psychological and sociological components.

Maleness or femaleness is determined at conception, when he inherits an X and a Y chromosome, or she inherits two X chromosomes. Another level of sexual identity comes from the control that hormones exert over the development of reproductive structures. Finally, both biological factors and social cues influence sexual feelings, including the strong sense of whether we are male or female.

Sexual Development

Gender differences do not become apparent until the ninth week of prenatal development. During the fifth week, all embryos develop two unspecialized gonads, which are organs that will develop as either testes or ovaries. Each such “indifferent” gonad forms near two sets of ducts that offer two developmental options. If one set of tubes, called the Müllerian ducts, continues to develop, they eventually form the sexual structures characteristic of a female. If the other set, the Wolffian ducts, persists, male sexual structures form.

The choice to follow a male or female developmental pathway occurs during the sixth week, depending upon the sex chromosome constitution and actions of certain genes. If a gene on the Y chromosome called *SRY* (for “sex-determining region of the Y”) is activated, hormones steer development along a male route. In the absence of *SRY* activation, a female develops (figure 6.1).

Femaleness was long considered a “default” option in human development, defined as the absence of maleness. Sex determination is more accurately described as a fate imposed on ambiguous precursor structures. Several genes besides *SRY* guide early development of reproductive structures. A mutation

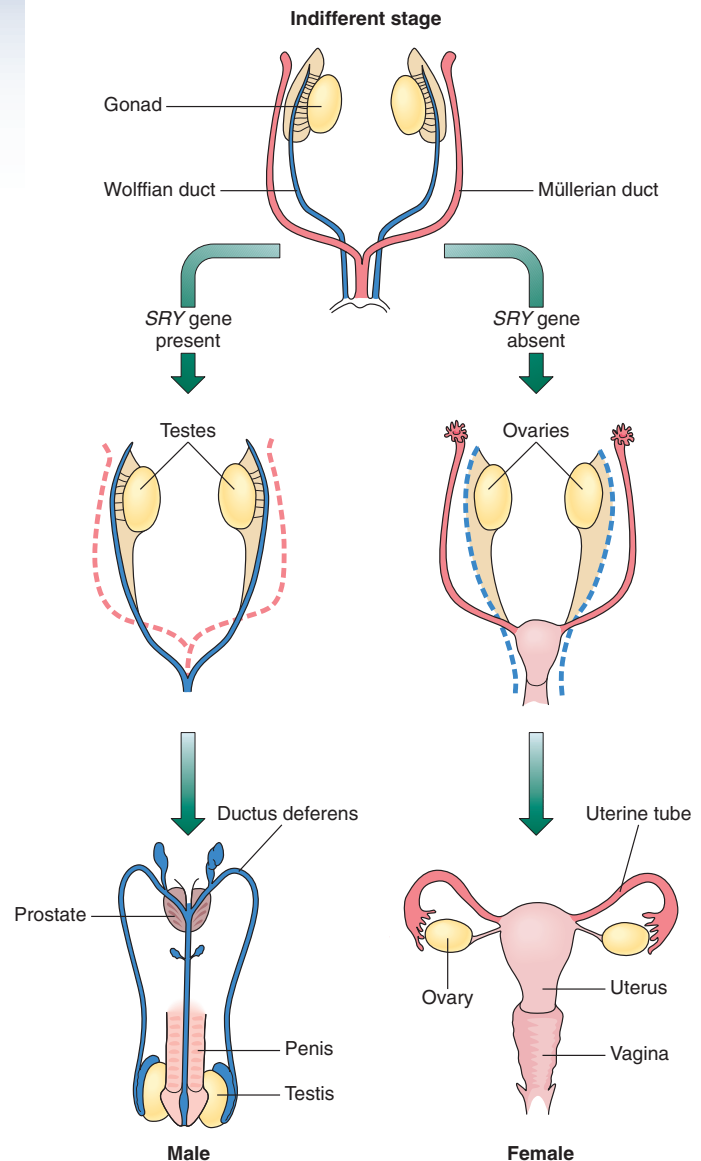
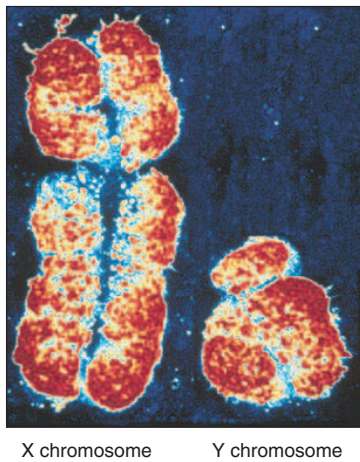


Figure 6.1 Male or female? The paired duct systems in the early human embryo may develop into male or female reproductive organs. The red tubes represent female structures and the blue tubes, male structures.

in a gene called *Wnt4*, for example, causes a female to have high levels of male sex hormones and lack a vaginal canal and uterus. Hence, the gene is essential for development and maturation as a female.

Sex Chromosomes

Human males and females have equal numbers of autosomes, but males have one X chromosome and one Y chromosome, and females have two X chromosomes (figure 6.2). The sex with two different sex chromosomes is called the **heterogametic**



X chromosome Y chromosome

Figure 6.2 Sex chromosomes. The human X and Y chromosomes are shown here in the replicated form because they were dividing when photographed.

sex, and the other, with two of the same sex chromosome, is the **homogametic sex**. In humans, males are heterogametic and females are homogametic. Some other species are different. In birds and snakes, for example, males are ZZ (homogametic) and females are ZW (heterogametic).

The sex chromosomes differ both in size and in gene content. The X chromosome in humans has more than 1,500 genes and is much larger than the Y chromosome, which has 231 protein-encoding genes. In meiosis in a male, the X and Y chromosomes act as if they are a pair of homologs. We introduce the Y chromosome here, then consider the X in section 6.2.

Identifying genes on the human Y chromosome has been difficult. Before the human genome sequence became available, researchers inferred the functions and locations of Y-linked genes by determining how men missing parts of the chromosome differ from normal. It was not possible to create linkage maps for the Y, as it was for the other chromosomes, because the Y does not cross over along all of its length.

Analysis of the genome sequence revealed why mapping the Y chromosome was so hard: It has a very unusual organization. In the 95 percent of the chromosome that harbors male-specific genes, many DNA segments are palindromes. In written languages, palindromes are sequences of letters that read the same in both directions—“Madam, I’m Adam,” for example. This symmetry in a DNA sequence, described by researchers as “a hall of mirrors,” destabilizes DNA replication. As a result, during meiosis, sections of a Y chromosome attract each other. This can loop out parts in between, which may account for many cases of male infertility caused by missing parts of the Y. Yet this “hall of mirrors” organization may also provide a way for the chromosome to recombine with itself, essentially sustaining its structure. Two researchers—one an XX, one an XY—take a lighthearted look at the

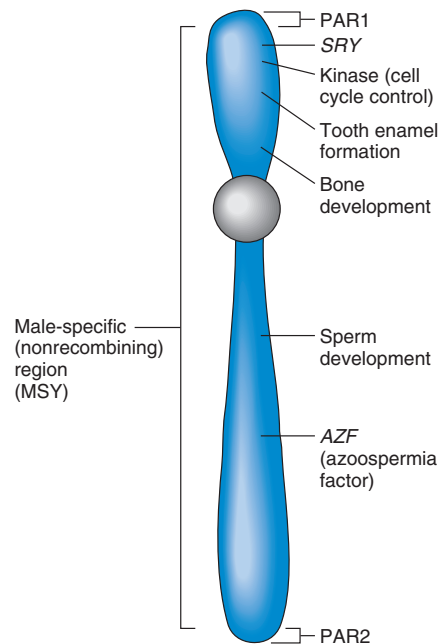


Figure 6.3 Anatomy of the Y chromosome. The Y chromosome has two pseudoautosomal regions (PAR1 and PAR2) and a large central area (MSY) that comprises about 95 percent of the chromosome. A few genes are indicated here. *SRY* determines sex. *AZF* encodes a protein essential to producing sperm; mutations in it cause infertility.

curious structure of the human Y chromosome in “In Their Own Words” on page 114.

The Y chromosome has a distinctive overall structure (**figure 6.3**) with a short arm and a long arm. At both tips of the Y chromosome are pseudoautosomal regions, termed PAR1 and PAR2. They comprise only 5 percent of the chromosome and include 63 pseudoautosomal genes. The term “pseudoautosomal” means that the DNA sequences have counterparts on the X chromosome and can cross over with them. The pseudoautosomal genes encode a variety of proteins that function in both sexes, participating in or controlling bone growth, cell division, immunity, signal transduction, the synthesis of hormones and receptors, fertility, and energy metabolism.

Most of the Y chromosome is the male-specific region, or MSY, that lies between the two pseudoautosomal regions. It consists of three classes of DNA sequences. About 10 to 15 percent of the MSY is X-transposed sequences that are 99 percent identical to counterparts on the X chromosome. Protein-encoding genes are scarce here. Another 20 percent of the MSY consists of X-degenerate DNA sequences, which are somewhat similar to X chromosome sequences and may be remnants of an ancient autosome that long ago gave rise to the X chromosome. The remainder of the MSY includes palindromes, called amplicons. The genes in the MSY include many repeats and specify protein segments that combine in different ways, which is one reason why counting

the number of protein-encoding genes on the Y chromosome has been difficult. Many of the genes in the MSY are essential to fertility, including *SRY*.

The Y chromosome was first seen under a light microscope in 1923, and researchers soon recognized its association with maleness. For many years, they sought to identify the gene or genes that determine sex. Important clues came from two very interesting types of people—men who have two X chromosomes (XX male syndrome), and women who have one X and one Y chromosome (XY female syndrome). A close look at their sex chromosomes revealed that the XX males actually had a small piece of a Y chromosome, and the XY females lacked a small part of the Y chromosome. The part of the Y chromosome present in the XX males was the same part that was missing in the XY females. This critical area accounted for half a percent of the Y chromosome. In 1990, researchers isolated and identified the *SRY* gene here.

The Phenotype Forms

The *SRY* gene encodes a very important type of protein called a **transcription factor**, which controls the expression of other genes. The *SRY* transcription factor stimulates male development by sending signals to the indifferent gonads. In response, sustentacular (also known as Sertoli) cells in the developing testis secrete anti-Müllerian hormone, which destroys potential female structures (uterus, uterine tubes, and upper vagina). At the same time, interstitial (also known as Leydig) cells in the testis secrete testosterone, which stimulates the development of male structures (the epididymides, ductus deferentia, seminal vesicles, and ejaculatory ducts). Some testosterone is also converted to dihydrotestosterone (DHT), which directs the development of the urethra, prostate gland, penis, and scrotum.

Because prenatal sexual development is a multistep process, genetic abnormalities can intervene at several different points (figure 6.4). The result may be an XY individual with a

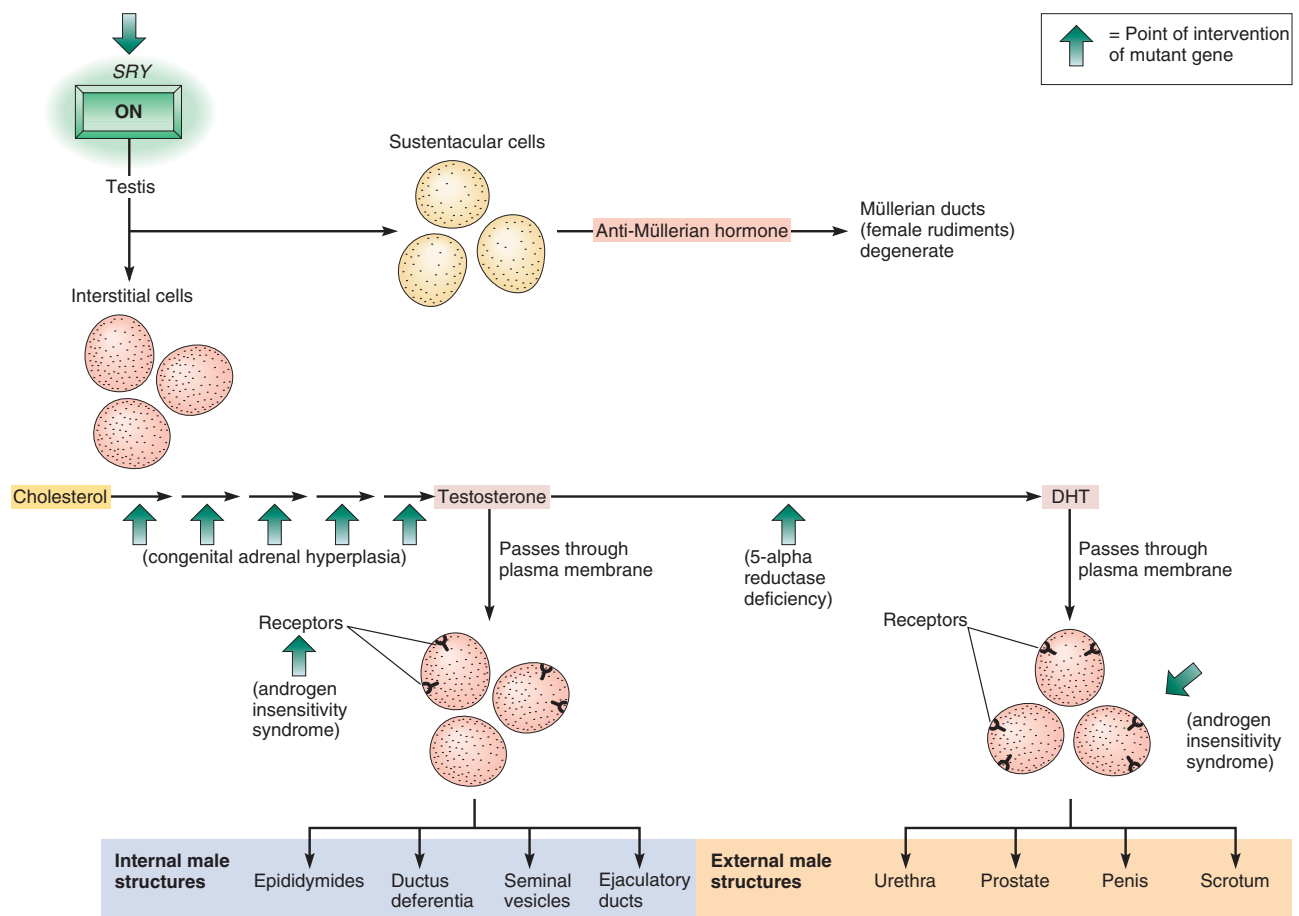


Figure 6.4 Pseudohermaphroditism. In normal male prenatal development, activation of a set of genes beginning with *SRY* stimulates sustentacular cells to produce hormones that lead to the destruction of female rudiments, and also stimulates interstitial cells to activate the biochemical pathway that produces testosterone and DHT. Testosterone and DHT promote the development of male structures. The green arrows indicate where mutations disrupt normal sexual development. (The ductus deferentia were formerly called vasa deferentia.)

In Their Own Words

The Y Wars

Researcher Jennifer Marshall-Graves predicts that the Y chromosome will “self-destruct” within the next 10 million years. Her comparison of Y chromosomes in a wide variety of mammals indicates that, gradually, important genes are being transferred to other chromosomes. David Page, who has led the mapping of the Y chromosome, has a more optimistic view. Each researcher spoke out, in jest, at two scientific conferences. Here is some of what they had to say:

The Rise and Fall of the Human Y Chromosome

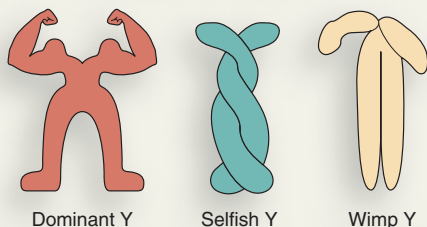
**Jennifer A. Marshall-Graves,
Australian National University**

The Y chromosome is unique in the human genome. It is small, gene-poor, prone to deletion and loss, variable among species, and useless. You can lack a Y and not be dead, just female. It is impossible to understand why this chromosome is so weird without understanding where it came from. It is a sad decline, and I predict its imminent loss.

The X is a decent sort of chromosome. It accounts for 5 percent of the genome, with about 1,500 perfectly normal genes. The Y is a pathetic little chromosome that has few genes interspersed with lots of junk. And those genes are a weird lot. They are particularly concerned with male sexual development, so they are rather specialized. There are a number of important genes, but some are quite bizarre and many inactive. The Y shares a lot of sequence with the X, and a lot of homology elsewhere, so the Y clearly diverged from the X.

There are several models of the Y (see figure). The dominant Y model of a macho Y reflects the fact that the Y contains the male-determining *SRY* gene. The selfish Y model predicts that the Y kidnapped genes from elsewhere. The wimp Y model says that the Y is just a relic of the once glorious X chromosome. This model was first proposed by biologist Susumo Ohno in 1967 in the theory that the X and Y originated as a pair of autosomes. Then the Y acquired

Models of the Human Y



Models of the human Y chromosome. Researcher Jennifer Marshall-Graves offers a tongue-in-cheek look at the Y chromosome, but her research findings are serious—the chromosome is shrinking.

the male-determining locus, and other genes that are required for spermatogenesis gathered nearby. This led to suppressed recombination in this region of the Y, which allowed all sorts of horrible genetic accidents to occur that could not be repaired. Mutations, deletions, and insertions accumulated until almost nothing was left, except bits at the top and bottom that still pair with the X. A few genes survived because they found a useful male-specific function, and many of these have made copies of themselves in a desperate race to stave off disappearing altogether.

The Y is degrading fast, losing genes at the rate of 5 per million years. I predict that it will be completely gone in 5 to 10 million years. Will we have males? The males in the audience can take comfort from the mole vole *Ellobius lutescens* (see photo). It has no Y, but it does have males and females. It has no *SRY*, no Y chromosome at all. Both sexes are XO. How do they do it? We don't know. Clearly another gene takes over and new sex genes start evolving. Will there be new sex chromosome evolution in humans? Maybe it will happen in different ways in different populations, and we will split into two species.

Rethinking The Rotting Y Chromosome

**David Page, Massachusetts Institute of Technology and
Howard Hughes Medical Institute Investigator**

The Y chromosome has had a public relations problem for a long time. For most of the last half of the past century, people thought that the Y chromosome was a junk heap. The genomic junkyard view was the classic model for sex chromosome evolution. We can now update that model.

Back 300 million years ago, when we were reptiles, we had no sex chromosomes, only ordinary autosomes. Shortly after our



Life without a Y? Males of all mammals, except two species of mole voles, have Y chromosomes. Birds and reptiles do not. The Y chromosome probably arose from an X chromosome about 310 million years ago. The X lost many genes and gained a few that set their carriers on the road to maleness. This animal is a Y-less male mole vole—it reproduces just fine.

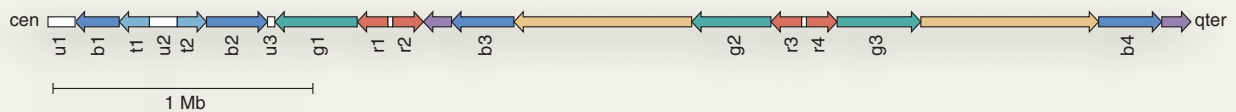
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ancestors parted company with the ancestors of birds, a mutation arose on one member of a pair of ordinary autosomes to give rise to *SRY*. The process of shutting down XY crossing over began, first in the vicinity of *SRY*, and then in an expanding region. Once a piece of the Y was no longer able to recombine with the X, its genes began to rot. The purpose of sex (recombination in meiosis) is not just to generate new gene combinations, but to allow genes to rid themselves of mildly deleterious mutations that accumulate. Y genes are not protected because they have lots of areas of no crossing over. Genes decayed, except for *SRY* and the tips. It wasn't a very flattering model for the Y.

When Jennifer Marshall-Graves and John Aitken wrote their article in *Nature* on the future of sex, that the Y would self-destruct in 10 million years, it truly frightened the people in my lab. We decided we needed to pick up the pace. When the popular press discovered the story of the impending death of the Y chromosome, they moved the date up to 5 million years from now.

Based on the sequencing of the Y, we've been able to rethink its evolution, and realized that the chromosome may have found a way around its seemingly inevitable problems. We looked closely at the male-specific region of the Y, reanalyzing sequences in a different way, chopped into smaller bits. And we found that each piece would find a match elsewhere on the Y. So segments on the Y are effectively functioning as alleles—30 percent have a perfect match elsewhere on the chromosome. These are not simple repeats, but highly complex sequences of tens to hundreds of kilobases. The region includes eight palindromes and one inverted repeat (see figure). We propose that there is intense recombination within the palindromes. And so the Y has two forms of productive recombination: conventional routine recombination of crossing over with the X at pseudoautosomal regions, and recombination within the Y. It's not that the Y doesn't recombine, it just does it its own way. The Y does copying that preserves its identity.



The Y chromosome is highly repetitive. A section of the Y chromosome that David Page studies, called *AZFc* (for azoospermia factor c), consists of DNA sequences that read the same in either direction, an organization that can lead to instability as well as provide a mechanism to generate new alleles. Other parts of the chromosome house similar repeats. Matching colors in this depiction represent identical sequences. Same-color arrows that point in opposite directions indicate inverted repeats.

block in the gene- and hormone-controlled elaboration of male structures so that the phenotype is female. For example, in androgen insensitivity syndrome (MIM 300068), caused by a mutation in a gene on the X chromosome, the absence of receptors for androgens (the male sex hormones testosterone and DHT) stops cells in early reproductive structures from receiving the signal to develop as male. The person looks female, but is XY.

Several terms are used to describe individuals whose genetic/chromosomal sex and physical structures, both internal and external, are not consistent with one gender. *Hermaphroditism* is an older and more general term for an individual with both male and female sexual structures. *Intersex* refers to individuals whose internal structures are inconsistent with external structures, or whose genitalia are ambiguous. It is the preferred term outside of medical circles. *Pseudohermaphroditism* refers to the presence of both types of structures but at different stages of life.

Living with pseudohermaphroditism can be confusing, and the quality of one's life depends on how accepting the society is of the condition. Consider 5-alpha reductase deficiency (MIM 264600), which is autosomal recessive. Affected individuals have a normal Y chromosome, a wild type *SRY* gene, and testes. Testosterone stimulates the Wolffian duct system to develop into the internal male reproductive tract, while anti-Müllerian hormone eliminates the female precursor structures, so the male anatomy makes sense on the inside. The outside

is a different story. When 5-alpha reductase, which normally catalyzes the reaction of testosterone to form DHT, is deficient, lack of DHT causes the fetus to develop externally as female—that is, without DHT, a penis cannot form. Changes begin to be noticed at puberty, when the adrenal glands, which sit atop the kidneys, start to produce testosterone. This now leads to masculinization. In this XY individual who thought she was female, the voice deepens, facial hair grows, and muscles begin to build an undeniably masculine physique. Instead of experiencing the expected breast development and menstruation, the clitoris enlarges into a penis. Usually sperm production is normal. XX individuals with 5-alpha reductase deficiency look female.

The degree to which pseudohermaphroditism disturbs the individual depends as much on society as it does on genetics. In the Dominican Republic in the 1970s, 22 young girls reached the age of puberty and began to transform into boys. They had a form of 5-alpha reductase deficiency that was fairly common due to consanguinity (relatives having children with relatives). The parents were happy that they had had sons after all, and so these special adolescents were given their own gender name—guededoces, for “penis at age 12.” They were fully accepted as whatever they wanted to be. Sadly this isn't always the case. A very realistic novel, *Middlesex*, tells the story of a young Greek-American man with this condition who grew up as a female.

In a more common form of pseudohermaphroditism, congenital adrenal hyperplasia (CAH) due to 21-hydroxylase deficiency (MIM 201910), an enzyme block causes testosterone and DHT to accumulate. It is autosomal recessive, and both males and females are affected. The higher levels of androgens cause precocious puberty in males or male secondary sex characteristics to develop in females. Boys may enter puberty as early as 3 years old, with well-developed musculature, small testes, and an enlarged penis. At birth, girls may have a swollen clitoris that appears to be a small penis. They are female internally, but as they reach puberty, the voice deepens, facial hair grows, and menstruation does not occur.

Prenatal tests that detect chromosomal sex have changed the way that pseudohermaphroditism is diagnosed. Before these tests were available, the condition was detected only after puberty, when masculinization occurred in a person who looked female. Today, pseudohermaphroditism is suspected when a prenatal chromosome check reveals an X and a Y chromosome, but the newborn is a phenotypic girl.

Transgender is a poorly understood condition related to sexual identity. A transgendered individual has the phenotype and sex chromosomes of one gender, but identifies extremely strongly with the opposite gender. It is a much more profound condition than transvestitism, which refers to a male who prefers women's clothing. The genetic or physical basis of transgender is not known. Some affected individuals undergo surgery so that their physical selves match the gender that they feel certain they are.

Is Homosexuality Inherited?

No one really knows why we have feelings of belonging to one gender or the other, or of being attracted to the same or opposite gender, but these feelings are intense. In homosexuality, a person's phenotype and genotype are consistent, and physical attraction is toward members of the same sex. Homosexuality is seen in all human cultures and has been observed for thousands of years. It has been documented in more than 500 animal species.

Evidence suggests that homosexuality reflects complex input from both genes and the environment. The genetic influence has been suspected for a long time on the basis of the strong feelings that homosexual individuals have as young children, well before they know of the existence or meaning of the term. Other evidence comes from identical twins, who are more likely to both be homosexual than are both members of fraternal twin pairs, suggesting a genetic component.

Experiments in the 1990s identified genetic markers on the X chromosome that tended to be identical among pairs of homosexual brothers, but not other pairs. This finding led to the idea that a single

gene, or a few genes, dictates sexual preference. These results could not be confirmed. Further studies on twins indicated what many have long suspected—the roots of homosexuality are not simple.

Twin studies compare a trait between identical and fraternal twin pairs, to estimate the rough proportion of a trait that can be attributed to genes. Chapter 7 discusses this approach further. Such a study done on all of the adult twins in Sweden found that in males, genetics contributes about 35 percent to homosexuality, whereas among females the genetic contribution is about 18 percent. Clearly, homosexuality reflects the input of many genes and environmental factors, and may in fact arise in a variety of ways.

Table 6.1 summarizes the several components of sexual identity.

Sex Ratio

Mendel's law of segregation predicts that populations should have approximately equal numbers of male and female newborns. That is, male meiosis should yield equal numbers of X-bearing and Y-bearing sperm. After birth, societal and environmental factors may favor survival of one gender over the other.

The proportion of males to females in a human population is called the **sex ratio**. Sex ratio is calculated as the number of males divided by the number of females multiplied by 1,000, for people of a particular age. (Some organizations describe sex ratio based on 1.0.) A sex ratio of equal numbers of males and females would be designated 1,000. The sex ratio at conception is called the primary sex ratio. In the United States for the past six decades, newborn boys have slightly outnumbered newborn girls, with the primary sex ratio averaging 1,050. The slight excess of boys may reflect the fact that Y-bearing sperm weigh slightly less than X-bearing sperm, and so they may reach the oocyte faster.

Sex ratio at birth is termed secondary and at maturity is called tertiary. Sex ratio can change markedly with age,

Table 6.1 Sexual Identity

Level	Events	Timing
Chromosomal/genetic	XY = male XX = female	Fertilization
Gonadal sex	Undifferentiated structure begins to develop as testis or ovary	6 weeks after fertilization
Phenotypic sex	Development of external and internal reproductive structures continues as male or female in response to hormones	8 weeks after fertilization, puberty
Gender identity	Strong feelings of being male or female develop	From childhood, possibly earlier
Sexual orientation	Attraction to same or opposite sex	From childhood

reflecting medical conditions that affect the sexes differently, as well as environmental factors that affect one sex more than the other, such as participation in combat or engaging in other dangerous behaviors.

It is interesting to see what happens when a society intentionally alters the sex ratio. This has been done in India and China, where researchers have identified great numbers of “missing females.” In these societies, prenatal diagnostic techniques are used to identify XX fetuses. Termination of XX fetuses, underreporting of female births, and, rarely, selective infanticide of girl babies all have contributed to a very unnaturally skewed sex ratio favoring males.

In China, by the year 2020, 20 million men will find themselves without female partners as a long-term consequence of that nation’s “one-child policy.” It began in 1979, with financial incentives to control runaway population growth. If a couple had a second child, the government revoked benefits. Some families, wanting their only child to be a boy, failed to continue or report female pregnancies. The reasoning was societal: A son would care for his aging parents, but a daughter would care for her in-laws.

China’s one-child policy prevented hundreds of millions of births. The average number of births per woman fell from 5.4 in 1971 to 1.8 in 2001 (**figure 6.5**). By the turn of the century, 117 boys were being born for every 100 girls. But the policy has led to a society today in which children have few siblings, cousins, aunts, or uncles. Young women are now rare, and have, ironically, become valued once again. The Chinese government is now promoting a “Care for Girls” program, which awards housing subsidies and scholarships to families that have girls. Government officials hope that by the 2010 census, a more natural sex ratio will show signs of returning.

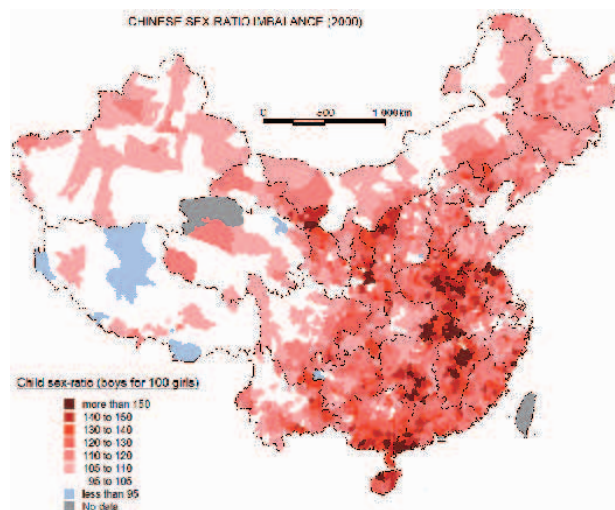


Figure 6.5 China’s one-child policy has led to a scarcity of women. In poor, rural areas, 94% of unwed young adults (aged 28 to 49) are men.

Today the same male bias that happened in China is occurring in India. Based on a survey of 1.1 million families, researchers found that the ratio of boys to girls was about equal when the first or second child was a boy, but if the first child and especially if the first two children were girls, then the secondary sex ratio fell to about 750 girls to every 1,000 boys. Families were using prenatal diagnosis to detect female pregnancies and were terminating about a fourth of them. Researchers estimate that India has about 100 million “missing females.”

At the other end of the human life cycle, sex ratio favors females in most populations. For people over the age of 65 in the United States, for example, the sex ratio is 720, meaning that there are 72 men for every 100 women. The ratio among older people is the result of disorders that are more likely to be fatal in males as well as behaviors that may shorten their life spans compared to women.

Key Concepts

1. The human female is XX and the male is XY.
2. The Y chromosome has few genes. It includes palindromes and sequences similar to sequences on the X chromosome.
3. Activation of *SRY* starts gene action that causes testes to develop in which sustentacular cells secrete anti-Müllerian hormone, which stops the development of female structures, and interstitial cells secrete testosterone, which stimulates development of male internal structures. Testosterone directs development of external structures. In several disorders, chromosomal, gonadal, and/or phenotypic sex are inconsistent.
4. Genes and the environment contribute to homosexuality.
5. Sex ratio is the number of males divided by the number of females multiplied by 1,000, for people of a particular age.

6.2 Traits Inherited on Sex Chromosomes

Genes carried on the Y chromosome are said to be **Y-linked**, and those on the X chromosome are **X-linked**. Y-linked traits are rare because the chromosome has few genes, and many have counterparts on the X chromosome. These traits are passed from male to male, because a female does not have a Y chromosome. No other Y-linked traits besides infertility (which obviously can’t be passed on) are yet clearly defined, although certain gene products have been identified. Claims that “hairy ears” is a Y-linked trait did not hold up—it turned out that families hid their affected female members!

Genes on the X chromosome have different patterns of expression in females and males, because a female has two X chromosomes and a male just one. In females, X-linked traits

are passed just like autosomal traits—that is, two copies are required for expression of a recessive allele, and one copy for a dominant allele. In males, however, a single copy of an X-linked allele causes expression of the trait or illness because there is no copy of the gene on a second X chromosome to mask the effect. A man inherits an X-linked trait only from his mother. The human male is considered **hemizygous** for X-linked traits, because he has only one set of X-linked genes.

Understanding how sex chromosomes are inherited is important in predicting phenotypes and genotypes in offspring. A male inherits his Y chromosome from his father and his X chromosome from his mother (**figure 6.6**). A female inherits one X chromosome from each parent. If a mother is heterozygous for a particular X-linked gene, her son or daughter has a 50 percent chance of inheriting either allele from her. X-linked traits are always passed on the X chromosome from mother to son or from either parent to daughter, but there is no male-to-male transmission of X-linked traits.

X-Linked Recessive Inheritance

An X-linked recessive trait is expressed in females if the causative allele is present in two copies. Many times, an X-linked trait passes from an unaffected heterozygous mother to an affected son. **Table 6.2** summarizes the transmission of an X-linked recessive trait.

If an X-linked condition is not lethal, a man may be healthy enough to transmit it to offspring. Consider the small family depicted in **figure 6.7**. A middle-aged man who had rough, brown, scaly skin did not realize his condition was

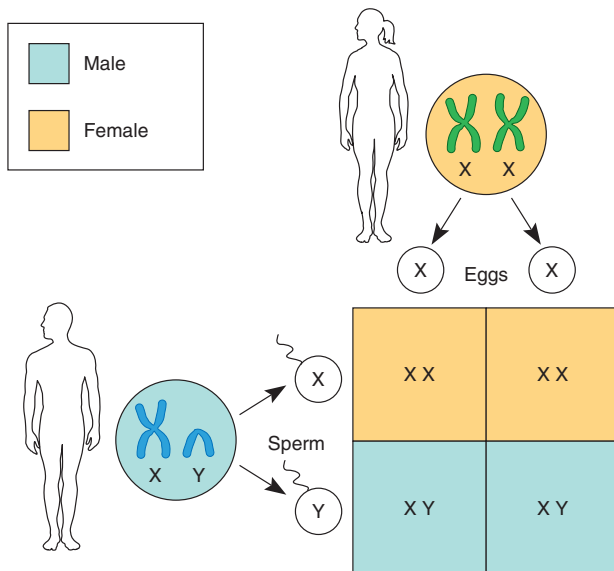


Figure 6.6 Sex determination in humans. An oocyte has a single X chromosome. A sperm cell has either an X or a Y chromosome. If a Y-bearing sperm cell with a functional *SRY* gene fertilizes an oocyte, the zygote is a male (XY). If an X-bearing sperm cell fertilizes an oocyte, then the zygote is a female (XX).

Table 6.2 Criteria for an X-Linked Recessive Trait

1. Always expressed in the male.
2. Expressed in a female homozygote but very rarely in a heterozygote.
3. Passed from heterozygote or homozygote mother to affected son.
4. Affected female has an affected father and a mother who is affected or a heterozygote.

inherited until his daughter had a son. By a year of age, the boy's skin resembled his grandfather's. In the condition, called ichthyosis (MIM 308100), an enzyme deficiency blocks removal of cholesterol from skin cells. The upper skin layer cannot peel off as it normally does, appearing brown and scaly. A test of the daughter's skin cells revealed that she produced half the normal amount of the enzyme, indicating that she was a carrier.

Colorblindness is another X-linked recessive trait that does not hamper the ability of a man to have children. About 8 percent of males of European ancestry are colorblind, as are 4 percent of males of African descent. Only 0.4 percent of females in both groups are colorblind. **Reading 6.1**, on page 119, takes a closer look at this interesting trait.

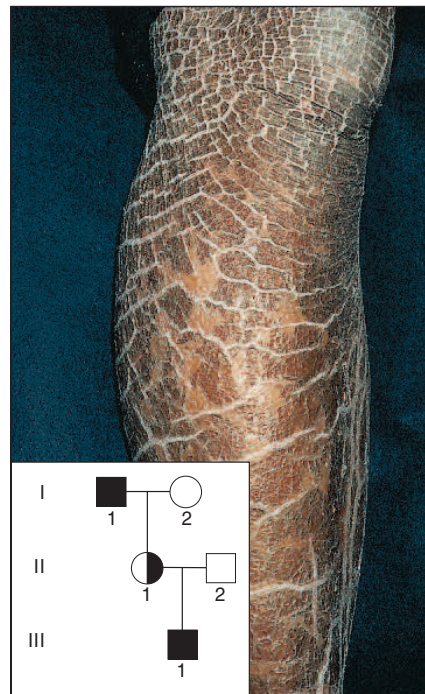


Figure 6.7 An X-linked recessive trait. Ichthyosis is transmitted as an X-linked recessive trait. A grandfather and grandson were affected in this family.

Reading 6.1

Colorblindness

English chemist John Dalton saw things differently from most people. Sealing wax that appeared red to other people was as green as a leaf to Dalton and his brother. Pink wildflowers were blue. The Dalton brothers had X-linked recessive colorblindness.

Curious about the cause of his colorblindness, Dalton asked his personal physician, Joseph Ransome, to dissect his eyes after he died. Ransome snipped off the back of one eye, removing the retina, where the cone cells that provide color vision are nestled among the more abundant rod cells that impart black-and-white vision. Because Ransome could see red and green normally when he peered through the back of his friend's eyeball, he concluded that it was not an abnormal filter in front of the eye that altered color vision. He stored the eyes in dry air, enabling researchers at the London Institute of Ophthalmology to analyze DNA in Dalton's eyeballs in 1994. Dalton's remaining retina lacked one of the three types of photopigments that enable cone cells to capture certain wavelengths of light.

Color Vision Basics

Cone cells are of three types, defined by the presence of any of three types of photopigments. An object appears colored because it reflects certain wavelengths of light, and each cone type captures a particular range of wavelengths with its photopigment. The brain then interprets the incoming information as a visual perception, much as an artist mixes the three primary colors to create many hues and shadings.

Each photopigment has a vitamin A-derived portion called retinal and a protein portion called an opsin. The three types of opsins correspond to short, middle, and long wavelengths of light. Mutations in opsin genes cause three different types of colorblindness.

A gene on chromosome 7 encodes shortwave opsins, and mutations in it produce the rare autosomal "blue" form of colorblindness (MIM 190900). Dalton had deuteranopia (green colorblindness), which means his eyes lacked the middle-wavelength opsin. In the third type, protanopia (red colorblindness), long-wavelength opsin is absent. Deuteranopia (MIM 303800) and protanopia (MIM 303900) are X-linked.

A Molecular View

Jeremy Nathans of Johns Hopkins University is another researcher who has personally contributed to our understanding of color vision. First, he used a cow version of a protein called rhodopsin that provides black-and-white vision to identify the human counterpart of the rhodopsin gene. Hypothesizing that the DNA sequence in the rhodopsin gene would be similar to that in the three opsin genes, and therefore able to bind to them, Nathans used the human rhodopsin gene as a "probe" to search his own DNA for genes with similar sequences. He found three. One was on chromosome 7, the other two on the X chromosome.

Although Nathans can see colors, his opsin genes are not

entirely normal, which provided a big clue to how colorblindness arises and why it is so common. On his X chromosome, Nathans has one red opsin gene and two green genes, instead of the normal one of each. Because the red and green genes have similar sequences, Nathans reasoned, they can misalign during meiosis in the female (**figure 1**). The resulting oocytes would then have either two or none of one opsin gene type. An oocyte lacking either a red or a green opsin gene would, when fertilized by a Y-bearing sperm, give rise to a colorblind male.

People who are colorblind must get along in a multicolored world. To help them overcome the disadvantage of not seeing important color differences, computer algorithms can convert colored video pictures into shades they can see.

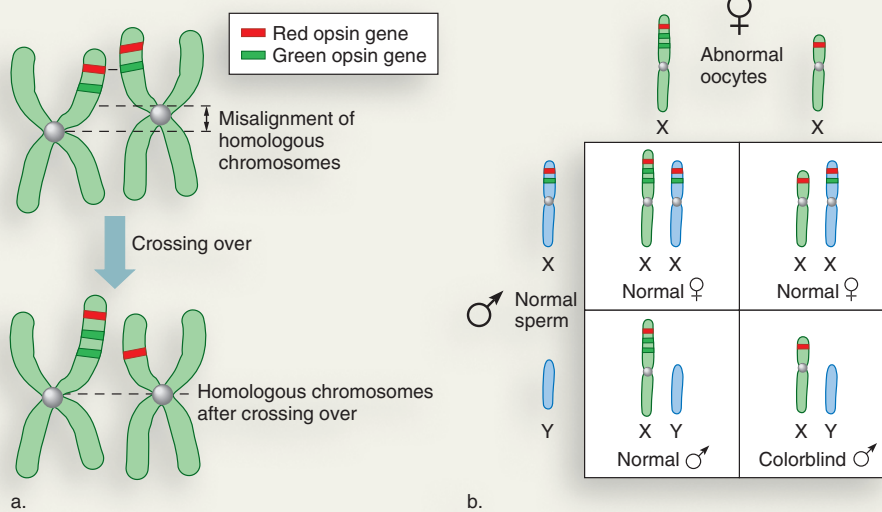


Figure 1 How colorblindness arises. **(a)** The sequence similarities among the opsin genes responsible for color vision may cause chromosomes to misalign during meiosis in the female. Offspring may inherit too many, or too few, opsin genes. A son inheriting an X chromosome missing an opsin gene would be colorblind. A daughter, unless her father is colorblind, would be a carrier. **(b)** A missing gene causes X-linked colorblindness.

Figure 6.8 shows part of an extensive pedigree for another X-linked recessive trait, the blood-clotting disorder hemophilia A (MIM 306700). Note the combination of pedigree symbols and a Punnett square to trace transmission of the trait. Dominant and recessive alleles are indicated by superscripts to the X and Y chromosomes. In the royal families of England, Germany, Spain, and Russia, the mutant allele arose in one of Queen Victoria's X chromosomes; it was either a new mutation or she inherited it. She passed it on through carrier daughters and one mildly affected son.

The transmission pattern of hemophilia A is consistent with the criteria for an X-linked recessive trait listed in

table 6.2. A daughter can inherit an X-linked recessive disorder or trait if her father is affected and her mother is a carrier, because the daughter inherits one affected X chromosome from each parent. Without a biochemical test, though, an unaffected woman would not know she is a carrier for an X-linked recessive trait unless she has an affected son.

A woman whose brother has hemophilia A has a 1 in 2 risk of being a carrier. Both her parents are healthy, but her mother must be a carrier because her brother is affected. Her risk is the chance that she has inherited the X chromosome bearing the hemophilia allele from her mother. The chance of the woman conceiving a son is 1 in 2, and of that son inheriting hemophilia is 1 in 2. Using the product rule, the risk that she

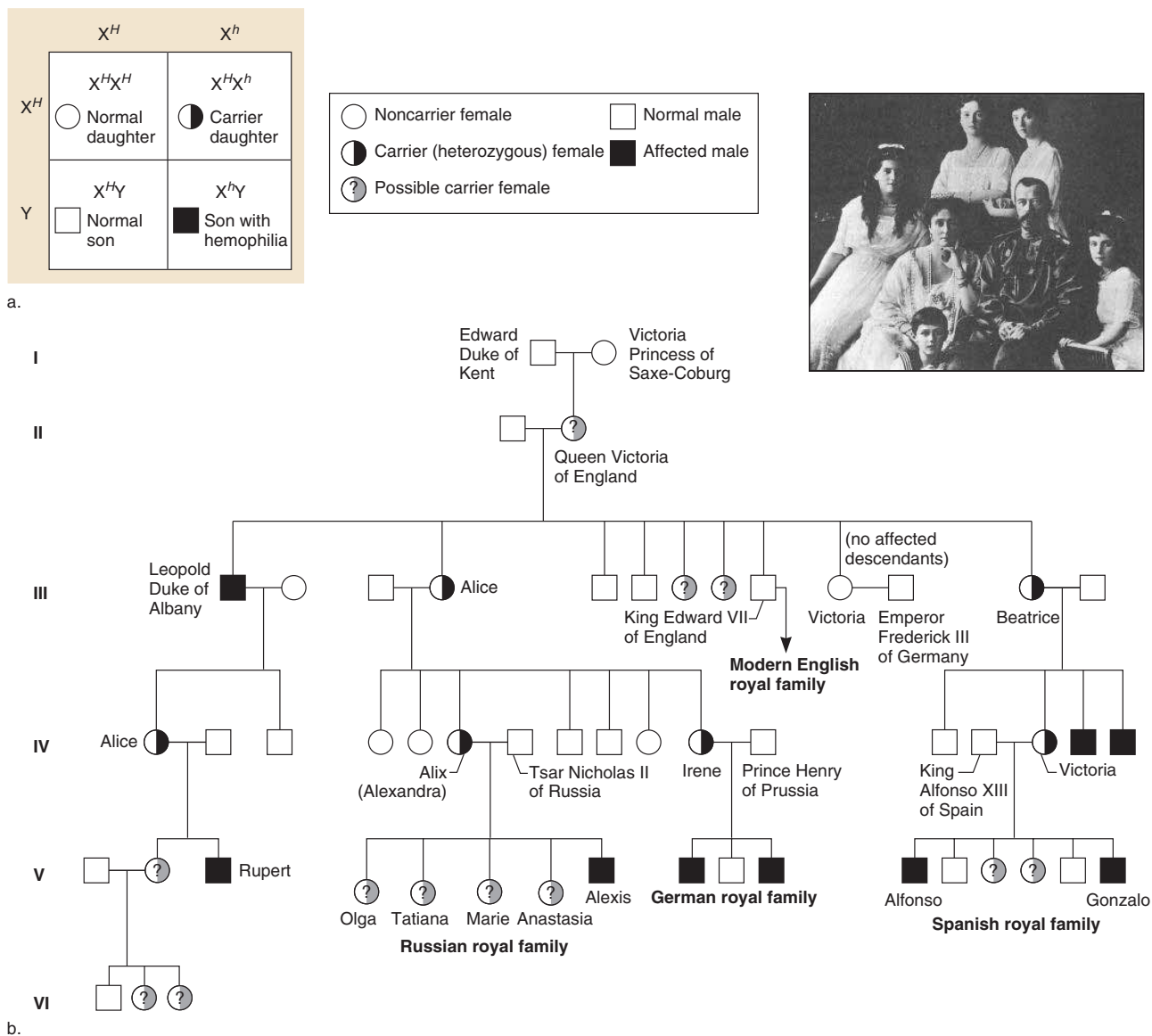


Figure 6.8 Hemophilia (a) This X-linked recessive disease usually passes from a heterozygous woman (designated $X^H X^h$, where h is the hemophilia-causing allele) to heterozygous daughters or hemizygous sons. The father is normal. (b) The disorder has appeared in the royal families of England, Germany, Spain, and Russia. The modern royal family in England does not carry hemophilia.

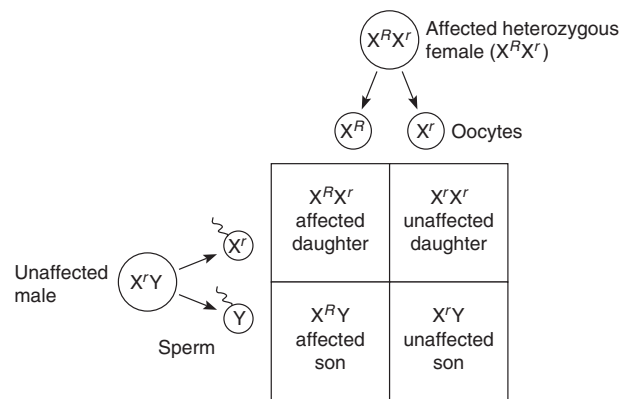
Table 6.3 Criteria for an X-Linked Dominant Trait

1. Expressed in female in one copy.
2. Much more severe effects in males.
3. High rates of miscarriage due to early lethality in males.
4. Passed from male to all daughters but to no sons.

is a carrier and will have a son with hemophilia, out of all the possible children she can conceive, is $1/2 \times 1/2 \times 1/2$, or $1/8$.

X-Linked Dominant Inheritance

Dominant X-linked conditions and traits are rare. Again, gene expression differs between the sexes (**table 6.3**). A female who inherits a dominant X-linked allele has the associated trait or illness, but a male who inherits the allele is usually more severely affected because he has no other allele to mask its effect. The children of a normal man and a woman with a dominant, disease-causing allele on the X chromosome face the risks summarized in **figure 6.9**.



a.



b.

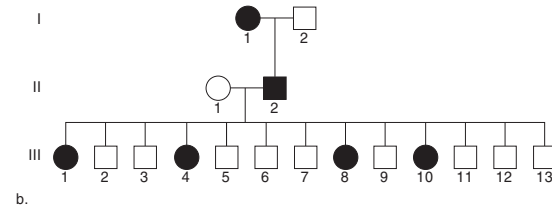
Figure 6.9 X-linked dominant inheritance. (a) A female who has an X-linked dominant trait has a 1 in 2 probability of passing it to her offspring, male or female. Males are generally more severely affected than females. (b) Note the characteristic patchy pigmentation on the leg of a girl who has incontinentia pigmenti.

An example of an X-linked dominant condition is incontinentia pigmenti (IP) (MIM 308300). In affected females, swirls of skin pigment arise when melanin penetrates the deeper skin layers. A newborn girl with IP has yellow, pus-filled vesicles on her limbs that come and go over the first few weeks. Then the lesions become warty and eventually give way to brown splotches that may remain for life, although they fade with time. Males with the condition are so severely affected that they do not survive to be born. This is why women with the disorder have a miscarriage rate of about 25 percent. The gene that causes IP (called *NEMO*) activates genes that carry out the immune response and apoptosis in tissues that derive from ectoderm, such as skin, hair, nails, eyes, and the brain.

Another X-linked dominant condition, congenital generalized hypertrichosis (MIM 307150), produces many extra hair follicles, and hence denser and more abundant upper body hair (**figure 6.10**). Hair growth is milder and patchier in females because of hormonal differences and the presence of a second X chromosome. Figure 6.10b shows part of a pedigree of a large Mexican family with 19 members who have this X-linked dominant condition. The affected male in the pedigree passed the trait to all four daughters, but to none of his nine sons. Because sons inherit the X chromosome from their



a.



b.

Figure 6.10 An X-linked dominant condition. (a) This 6-year-old child has congenital generalized hypertrichosis. (b) In this partial pedigree of a large Mexican family, the affected male in the second generation passed the condition to all of his daughters and none of his sons. This is because he transmits his X chromosome only to females.

mother, and only the Y from their father, they could not have inherited the hairiness from their father.

+ Solving a Problem

X-Linked Inheritance

Mendel's first law (segregation) applies to genes on the X chromosome. Therefore, the same logic is used to solve problems as to trace traits transmitted on autosomes, with the added step of considering the X and Y chromosomes in Punnett squares. Follow these steps:

- Look at the pattern of inheritance. Different frequencies of affected males and females in each generation may suggest X linkage.
For an X-linked recessive trait:
 - An affected male has a carrier mother.
 - An unaffected female with an affected brother has a 50 percent (1 in 2) chance of being a carrier.
 - An affected female has a carrier or affected mother *and* an affected father.
 - A carrier (female) has a carrier mother *or* an affected father.
 For an X-linked dominant trait:
 - There may be no affected males, because they die early.
 - An affected female has an affected mother.
- Draw the pedigree.
- List all genotypes and phenotypes and their probabilities.
- Assign genotypes and phenotypes to the parents. Consider clues in the phenotypes of relatives.
- Determine how alleles separate into gametes for the genes of interest on the X and Y chromosomes.
- Unite the gametes in a Punnett square.
- Determine the phenotypic and genotypic ratios for the F_1 generation.
- To predict further generations, use the genotypes of the F_1 and repeat steps 4 through 6.

Consider Kallmann syndrome (MIM 308700), which causes very poor or absent sense of smell and small testes or ovaries. It is X-linked recessive. Tanisha does not have Kallmann syndrome, but her brother Jamal and her maternal cousin Malcolm (her mother's sister's child) have it. Tanisha's and Malcolm's parents are unaffected, as is Tanisha's husband Sam. Tanisha and Sam wish to know the risk that a son would inherit the condition. Sam has no affected relatives.

Solution

- Mode of inheritance: The trait is X-linked recessive because males are affected through carrier mothers.

- K = wild type k = Kallmann syndrome

Genotypes	Phenotypes
X^KX^K , X^KX^k , X^KY	normal
X^kX^k , X^kY	affected

Individual	Genotype	Phenotype	Probability
Tanisha	X^KX^k or X^KX^K	normal (carrier)	50% each
Jamal	X^kY	affected	100%
Malcolm	X^kY	affected	100%
Sam	X^KY	normal	100%

- Tanisha's gametes

if she is a carrier:	X^K	X^k
Sam's gametes:	X^K	Y

- Punnett square

	X^K	X^k
X^K	X^KX^K	X^KX^k
Y	X^KY	X^kY

- Interpretation: If Tanisha is a carrier, the probability that their son will have Kallmann syndrome is 50 percent, or 1 in 2. (This is a conditional probability. The chance that any son will have the condition is actually 1 in 4, because Tanisha also has a 50 percent chance of being genotype X^KX^K and therefore not a carrier.)

Key Concepts

- Y-linked traits are passed on the Y chromosome, and X-linked traits on the X.
- Because a male is hemizygous, he expresses all the genes on his X chromosome, whereas a female expresses recessive alleles on the X chromosome only if she is homozygous recessive.
- X-linked recessive traits have a 50 percent probability of passing from carrier mothers to sons.
- X-linked dominant conditions are expressed in both males and females but are more severe in males.
- Mendel's first law can be used to solve problems involving X-linked genes.

6.3 Sex-Limited and Sex-Influenced Traits

An X-linked recessive trait generally is more prevalent in males than females. Other situations, however, can affect gene expression in the sexes differently.

Sex-Limited Traits

A **sex-limited trait** affects a structure or function of the body that is present in only males or only females. Such a gene may be X-linked or autosomal.

Understanding sex-limited inheritance is important in animal breeding. For example, a New Zealand cow named Marge, who has a mutation that makes her milk very low in saturated fat, is founding a commercial herd. Males play their part by transmitting the mutation, even though they do not make milk. In humans, beard growth is sex-limited. A woman does not grow a beard because she does not manufacture the hormones required for facial hair growth. She can, however, pass to her sons the genes specifying heavy beard growth.

An inherited medical condition that arises during pregnancy is obviously sex-limited, but the male genome contributes to the development of supportive structures, such as the placenta. This is the case for preeclampsia, a sudden rise in blood pressure late in pregnancy. It kills 50,000 women worldwide each year. A study of 1.7 million pregnancies in Norway found that if a man's first wife had preeclampsia, his second wife had double the risk of developing the condition, too. Another study found that women whose mothers-in-law developed preeclampsia when pregnant with the women's husbands had approximately twice the rate of developing the condition themselves. Perhaps a gene from the father affects the placenta in a way that elevates the pregnant woman's blood pressure.

Sex-Influenced Traits

In a **sex-influenced trait**, an allele is dominant in one sex but recessive in the other. Such a gene may be X-linked or autosomal. The difference in expression can be caused by hormonal differences between the sexes. For example, an autosomal gene for hair growth pattern has two alleles, one that produces hair all over the head and another that causes pattern baldness. The baldness allele is dominant in males but recessive in females, which is why more men than women are bald. A heterozygous male is bald, but a heterozygous female is not. A bald woman is homozygous recessive. Even a bald woman tends to have some wisps of hair, whereas an affected male may be completely hairless on the top of his head.

Key Concepts

1. A sex-limited trait affects body parts or functions present in only one gender.
2. A sex-influenced allele is dominant in one sex but recessive in the other.

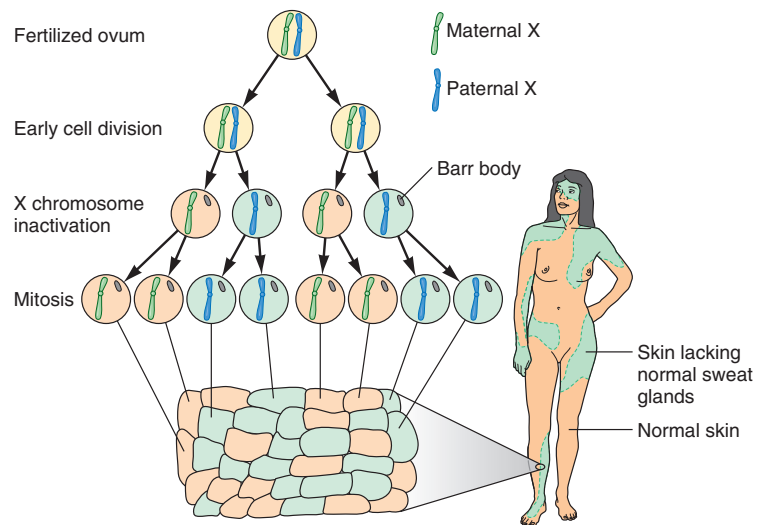


Figure 6.11 X inactivation. A female is a mosaic for expression of genes on the X chromosome because of the random inactivation of either the maternal or paternal X in each cell early in prenatal development. In anhidrotic ectodermal dysplasia, a woman has patches of skin that lack sweat glands and hair. (Colors distinguish cells with the inactivated X, not to depict skin color.)

6.4 X Inactivation

Females have two alleles for every gene on the X chromosomes, whereas males have only one. In mammals, a mechanism called **X inactivation** balances this apparent inequality in the expression of genes on the X chromosome.

Equaling Out the Sexes

By the time a female embryo consists of 8 cells, about 75 percent of the genes on one X chromosome in each cell are inactivated, and the remaining 25 percent are expressed to different degrees in different women. Which X chromosome is mostly turned off in each cell—the one inherited from the mother or the one from the father—is usually random. As a result, a female mammal expresses the X chromosome genes inherited from her father in some cells and those from her mother in others. She is, therefore, a mosaic for expression of most genes on the X chromosome (**figure 6.11**).

By studying rare human females who have lost a small part of one X chromosome, researchers identified a specific region, the X inactivation center, that shuts off much of the chromosome. Genes in the PARs and some other genes escape inactivation. A gene called *XIST* controls X inactivation. It encodes an RNA that binds to a specific site on the same (inactivated) X chromosome. From this point out to the chromosome tip, the X chromosome is inactivated.

Once an X chromosome is inactivated in one cell, all its daughter cells have the same X chromosome inactivated. Because the inactivation occurs early in development, the adult female has patches of tissue that differ in their expression of

X-linked genes. With each cell in her body having only one active X chromosome, she is roughly equivalent to the male in terms of gene expression.

X inactivation can alter the phenotype (gene expression), but not the genotype. It is not permanent, and is reversed in germline cells destined to become oocytes. Therefore, a fertilized ovum does not have an inactivated X chromosome.

X inactivation is an example of an **epigenetic** change—one that is passed from one cell generation to the next but that does not alter the DNA base sequence. We can observe X inactivation at the cellular level because the turned-off X chromosome absorbs a stain much faster than the active X. This differential staining occurs because inactivated DNA has chemical methyl (CH_3) groups that prevent it from being transcribed into RNA and also enable it to absorb stain.

X inactivation can be used to check the sex of an individual. The nucleus of a cell of a female, during interphase, has one dark-staining X chromosome called a **Barr body**. A cell from a male has no Barr body because his one X chromosome remains active.

In 1961, English geneticist Mary Lyon proposed that the Barr body is the inactivated X chromosome and that it is turned off in early development. Checking for Barr bodies has been done in the Olympics to identify athletes competing as the wrong gender.

Effect on the Phenotype

The consequence of X inactivation on the phenotype can be interesting. For homozygous X-linked genotypes, X inactivation has no effect. No matter which X chromosome is turned off, the same allele is left to be expressed. For heterozygotes, however, X inactivation leads to expression of one allele or the other. This doesn't affect health if enough cells express the functional gene product. However, some traits reveal the X inactivation. The swirls of skin color in incontinentia pigmenti (IP) patients reflect patterns of X inactivation in skin cells (see fig. 6.9b). Where the normal allele for melanin pigment is shut off, pale swirls develop. Where pigment is produced, brown swirls result.

A female who is heterozygous for an X-linked recessive gene can express the associated condition if the normal allele is inactivated in the tissues that the illness affects. Consider a carrier of hemophilia A. If the X chromosome carrying the normal allele for the clotting factor is turned off in the liver, then the woman's blood will clot slowly enough to cause mild hemophilia. (Luckily for her, slowed clotting time also greatly reduces her risk of cardiovascular disease caused by blood clots blocking circulation.) A carrier of an X-linked trait who expresses the phenotype is called a **manifesting heterozygote**.

Whether or not a manifesting heterozygote results from X inactivation depends upon how adept cells are at sharing. Consider two lysosomal storage disorders, which are deficiencies of specific enzymes that normally dismantle cellular debris in lysosomes. In Hunter syndrome (MIM 309900, also called mucopolysaccharidosis II), cells that make the enzyme readily send it to neighboring cells that do not, essentially correcting the defect in cells that can't make the enzyme. Carriers of Hunter syndrome do not have symptoms because cells get enough

enzyme. Affected boys are deaf, mentally retarded, have dwarfism and abnormal facial features, heart damage, and enlarged liver and spleen. In contrast, in Fabry disease (MIM 301500, also called alpha-galactosidase A deficiency), the enzyme is not easily released from cells, so a female who is a heterozygote may have cells in the affected organs that lack the enzyme. She may develop mild symptoms of this disorder that causes skin lesions, abdominal pain, and kidney failure in boys.

A familiar example of X inactivation is the coat colors of tortoiseshell and calico cats. An X-linked gene confers brownish-black (dominant) or yellowish-orange (recessive) color. A female cat heterozygous for this gene has patches of each color, forming a tortoiseshell pattern that reflects different cells expressing either of the two alleles (**figure 6.12**). The earlier the X inactivation, the larger the patches, because more cell divisions can occur after the event, producing more daughter cells. White patches may form due to epistasis by an autosomal gene that shuts off pigment synthesis. A cat with colored patches against such a white background is a calico. Tortoiseshell and calico cats are nearly always female. A male can have these coat patterns only if he inherits an extra X chromosome.

In humans, X inactivation can be used to identify carriers of some X-linked disorders. This is the case for Lesch-Nyhan syndrome (MIM 300322), in which an affected boy has cerebral palsy, bites his fingers, shoulders, and lips to the



Figure 6.12 Visualizing X inactivation. X inactivation is obvious in a calico cat. X inactivation is rarely observable in humans because most cells do not remain together during development, as a cat's skin cells do.

point of mutilation, is mentally retarded, and passes painful urinary stones. Mutation results in defective or absent HGPRT, an enzyme. A woman who carries Lesch-Nyhan syndrome can be detected when hairs from widely separated parts of her head are tested for HGPRT. (Hair is used for the test because it is accessible and produces the enzyme.) If some hairs contain HGPRT but others do not, she is a carrier. The hair cells

that lack the enzyme have turned off the X chromosome that carries the normal allele; the hair cells that manufacture the normal enzyme have turned off the X chromosome that carries the disease-causing allele. The woman is healthy because her brain has enough HGPRT, but each son has a 50 percent chance of inheriting the disease. **Reading 6.2** discusses another syndrome affected by X inactivation.

Reading 6.2

Rett Syndrome—A Curious Inheritance Pattern

Before the age of the Internet, identifying and describing a new syndrome could take years, or even decades. This was the case for Rett syndrome, a neurological condition that affects nearly always females.

In 1954, Austrian pediatrician Andreas Rett and his nurse noticed that their practice included eight little girls who did strange things with their hands—uncontrollably (**figure 1**). They'd tap objects, clap, repeatedly put their hands in their mouths, and most commonly, wring their hands. The girls shared other symptoms. All had been developing normally but then gradually lost muscle tone. Growth of their heads slowed. As time went on, seizures began, they lost the ability to speak, and the girls became completely disabled (**table 1**).

Dr. Rett filmed the girls and went around Europe looking for other cases. Meanwhile, other pediatricians were noting the symptoms in their patients, independently. Although Dr. Rett published his observations in European journals, they did not attract the attention of the medical mainstream until 1983, when a Swedish researcher published in the *Annals of Neurology* and, finally, others noticed. He named the condition Rett syndrome.

In 1999, Ruthie Amir, at the Baylor College of Medicine in Texas, discovered the gene behind the disorder—*MECP2*, for methyl-CpG-binding protein 2 (MIM 312750), on the X chromosome. The syndrome is pleiotropic, causing symptoms in several organ systems, because the gene adds methyl groups to other genes, silencing them.

Rett syndrome is dominant.

In 99 percent of cases, it is not passed from parent to child but arises anew. This is not surprising, because affected parents of either sex would be too ill to have children. The disease arises in either of two ways: The gene may mutate in an X-bearing sperm cell; alternatively, an affected girl's mother might indeed have the syndrome, but due to skewed X inactivation, the X chromosome carrying the mutation is turned off in most brain cells. The mother's case might be so mild as to be undetected—hence, her passing it to her daughter would appear as a sporadic (noninherited) case.



Figure 1 Rett syndrome affects girls. One sign of Rett syndrome is holding and wringing the hands.

Table 1 Stages and Symptoms of Rett Syndrome

Stage	Onset	Symptoms
I	6–18 months	Minor slowing of development; loss of eye contact and interest in objects.
II	1–4 years	Loss of ability to speak; autistic behavior; breathing irregularities; unsteady gait; head growth slows; mental retardation.
III	2–10 years	Seizures and loss of motor skills, but behavior may improve and autistic features fade; this stage can last a lifetime.
IV	adolescence	Severe motor problems, including rigidity, weakness, adulthood and spasticity; walking impaired, but cognition and communication skills remain.

Subtle Effects of X Inactivation

Theoretically, X inactivation evens out the sexes for expression of X-linked genes. In actuality, however, a female may *not* be equivalent, in gene expression, to a male because she has two cell populations, whereas a male has only one. One of a female's two cell populations has the X she inherited from her father active, and the other has the X chromosome she inherited from her mother active. For heterozygous X-linked genes, she would have some cells that manufacture the protein encoded by one allele, and some cells that produce the protein encoded by the other allele. Although most heterozygous genes have the alleles about equally represented, sometimes X inactivation can be skewed. That is, most cells express the X inherited from the same parent. This can happen if one of the X chromosomes includes an expressed allele that confers a greater rate of cell division than the different allele from the other parent, giving certain cells a survival advantage.

Another way that X inactivation makes a female different from a male is seen when the proteins encoded by different alleles interact. This can be beneficial or harmful. A beneficial example of dual expression of alleles occurs in certain types of monkeys in which an X-linked visual pigment gene has two alleles. Females who are homozygous for this gene and males have 2-color vision, but lucky female monkeys who are heterozygous for this gene enjoy 3-color vision.

A situation in which being a heterozygote for an X-linked gene is harmful is craniofrontonasal syndrome (MIM 304110) (figure 6.13). Males and homozygous females have asymmetrical facial features. However, heterozygous females have a much more severe phenotype, with very abnormal faces resulting from abnormal fusing of the skull bones. (It is highly unusual



Figure 6.13 Craniofrontonasal syndrome is more severe in females because of an unusual detrimental effect of expressing both alleles of an X-linked gene.

for the heterozygote to be more severely affected than the homozygous recessive individual.) An explanation is that the encoded protein is part of a signal transduction pathway that controls the bone fusion, and when two forms of that protein are made in the female heterozygote, the signal is disrupted in a way that blocks the cells that form the sutures of the skull from joining cleanly.

Key Concepts

1. In female mammals, X inactivation compensates for differences between males and females in the numbers of gene copies on the X chromosome.
2. Early in development, one X chromosome in each cell of the female is turned off.
3. The effects of X inactivation can be noticeable when heterozygous alleles are expressed in certain tissues.

6.5 Genomic Imprinting

In Mendel's pea experiments, it didn't matter whether a trait came from the male or female parent. For certain genes in mammals, however, parental origin does influence the phenotype. These genes are said to be imprinted. In **genomic imprinting**, methyl (CH_3) groups cover a gene or several linked genes and prevent them from being accessed to synthesize protein (figure 6.14).

For a particular imprinted gene, the copy inherited from either the father or the mother is always covered with methyls, even in different individuals. The result of this gene cloaking is that a disease may be more severe, or different, depending upon which parent transmitted the mutant allele. That is, a particular gene might function if it came from the father, but not if it came from the mother, or vice versa.



Figure 6.14 Methyl (CH_3) groups (red) "silence" certain genes.

Silencing the Contribution From One Parent

Imprinting is an epigenetic alteration. It is a layer of meaning stamped upon a gene without changing its DNA sequence. The imprinting pattern is passed from cell to cell in mitosis, but not from individual to individual through meiosis. When silenced DNA is replicated during mitosis, the pattern of blocked genes is exactly placed, or imprinted, on the new DNA, covering the same genes as in the parental DNA (figure 6.15). In this way, the “imprint” of inactivation is perpetuated, as if each such gene “remembers” which parent it came from. In meiosis, however, imprints are removed and reset. As oocyte and sperm form, the CH₃ groups shielding their imprinted genes are stripped away, and new patterns are set down, depending upon whether the fertilized ovum chromosomally is male (XY) or female (XX). In this way, women can have sons and men can have daughters without passing on their sex-specific parental imprints.

The function of genomic imprinting isn't well understood, but because many imprinted genes take part in early development, particularly of the brain, it may be a way to finely regulate the abundance of key proteins in the embryo. The fact that some genes lose their imprints after birth supports this idea of early importance. Also, imprinted genes are in clusters along a chromosome, and are controlled by other regions of DNA called imprinting centers. Perhaps one gene in a cluster is essential for early development, and the others become imprinted simply because they are nearby—a bystander effect.

Genomic imprinting has implications for understanding early human development. It suggests that for mammals, two opposite-sex parents are necessary to produce a healthy embryo and placenta. This apparent requirement for opposite-sex parents was discovered in the early 1980s, through experiments on early mouse embryos and examination of certain rare pregnancy problems in humans. Researchers created fertilized mouse ova that contained two male pronuclei or two female pronuclei, instead of one from each. Results were strange. When the fertilized ovum had two male genomes, a normal placenta developed, but the embryo was tiny and quickly stopped developing. A zygote with two female pronuclei, on the other hand, developed into an embryo, but the placenta was grossly abnormal. Therefore, the male genome controls placenta development, and the female genome, embryo development.

The mouse results were consistent with abnormalities of human development. When two sperm fertilize an oocyte

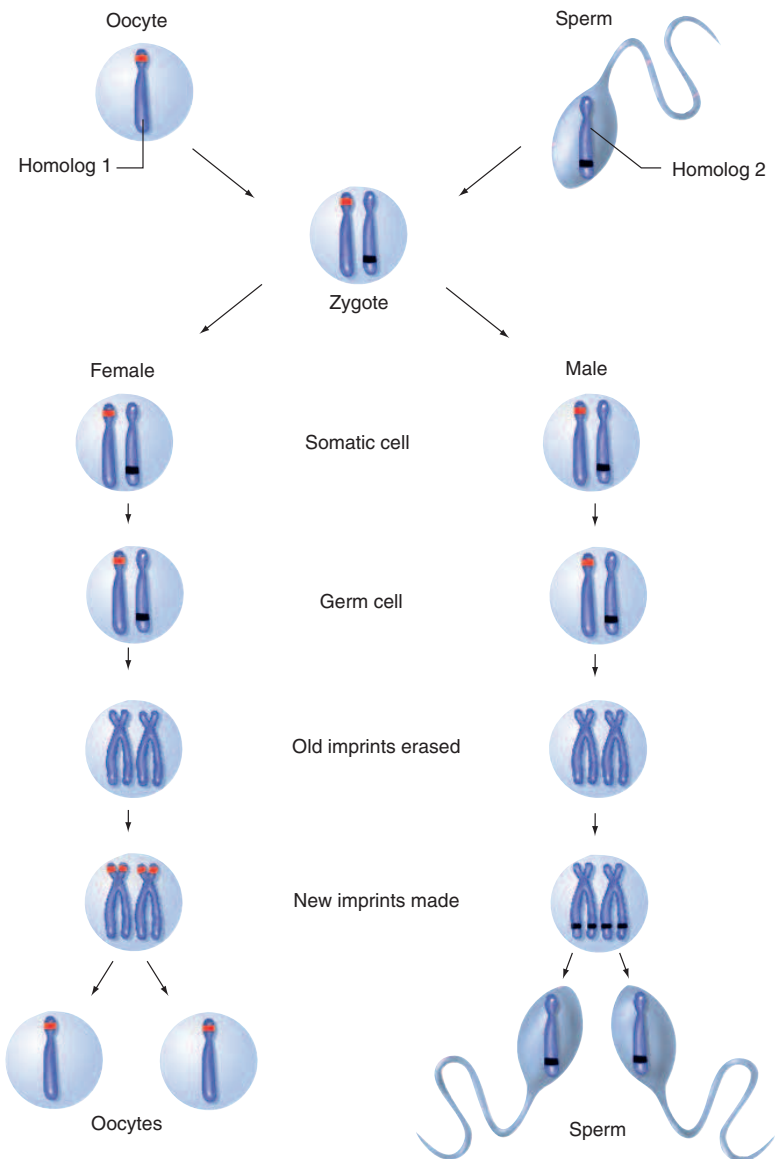


Figure 6.15 Genomic imprinting. Imprints are erased during meiosis, then reinstated according to the sex of the new individual.

and the female pronucleus degenerates, an abnormal growth of placenta-like tissue called a hydatidiform mole forms. If a fertilized ovum contains only two female genomes but no male genome, a mass of random differentiated tissue, called a teratoma, grows. A teratoma, which means “monster cancer,” may consist of a variety of tissues in a bizarre mix. With either a hydatidiform mole or a teratoma, no embryo results, although a pregnancy test may be positive, because the pregnancy hormone (hCG) may be produced.

Genomic imprinting can explain incomplete penetrance, in which an individual is known to have inherited a genotype associated with a particular phenotype, but has no signs of the

trait—such as a person with normal fingers whose parent and child have polydactyly. An imprinted gene that silences the dominant mutant allele could explain these cases: The predicted genotype is present, but the associated phenotype is not expressed.

Imprinting may be an important concern in assisted reproductive technologies that manipulate gametes to treat infertility. For example, the otherwise very rare Angelman syndrome (MIM 105830) and Beckwith-Wiedemann syndrome (MIM 130650) are more prevalent among the offspring of people who used *in vitro* fertilization and intracytoplasmic sperm injection (discussed in chapter 21) to become pregnant.

Imprinting Disorders in Humans

A genome-wide search for imprinted genes identified 156, at least 60 of which affect health when abnormally expressed. The effects of genomic imprinting are revealed only when an individual has one copy of a normally imprinted allele and the other, active allele is inactivated or deleted. The effects of genomic imprinting gone awry can be dramatic, such as two different syndromes that arise from small deletions in the same region of chromosome 15 (**figure 6.16**). A child with Prader-Willi syndrome (MIM 176270) is small at birth and in infancy has difficulty gaining weight. Between ages 1 and 3, the child develops an obsession with eating. Unless the diet can be controlled, severe obesity results because another symptom is a very slow metabolism. Parents actually lock kitchen cabinets and refrigerators to keep their children from literally eating themselves to death by bursting digestive organs. The other condition, Angelman syndrome, causes autism and mental retardation, an extended tongue, large jaw, poor muscle coordination, and convulsions that make the arms flap. In many cases of Prader-Willi syndrome, only the mother's chromosome 15 region is expressed; the father's chromosome is deleted in that region. In Angelman syndrome, the reverse occurs: The father's gene (or genes) is expressed, and the mother's chromosome has the deletion.

Symptoms of Prader-Willi arise because several paternal genes that are not normally imprinted (that is, that are normally active) are missing. In Angelman syndrome, a normally active single maternal gene is deleted. This part of chromosome 15 is especially unstable because it includes highly repetitive DNA sequences, which bracket the genes that cause the symptoms.

Imprinting gone awry is associated with forms of diabetes mellitus, autism, Alzheimer disease, schizophrenia, and male homosexuality. It was the association of autism in some syndromes with expression of the father's genes that contributed to the imprinted brain hypothesis described in the chapter opening essay. Clues that indicate a condition is associated with genomic imprinting include increased severity depending on whether it is inherited from the father or mother and also a phenomenon called **uniparental disomy**. This term literally means “two bodies from one parent,” and refers to an offspring

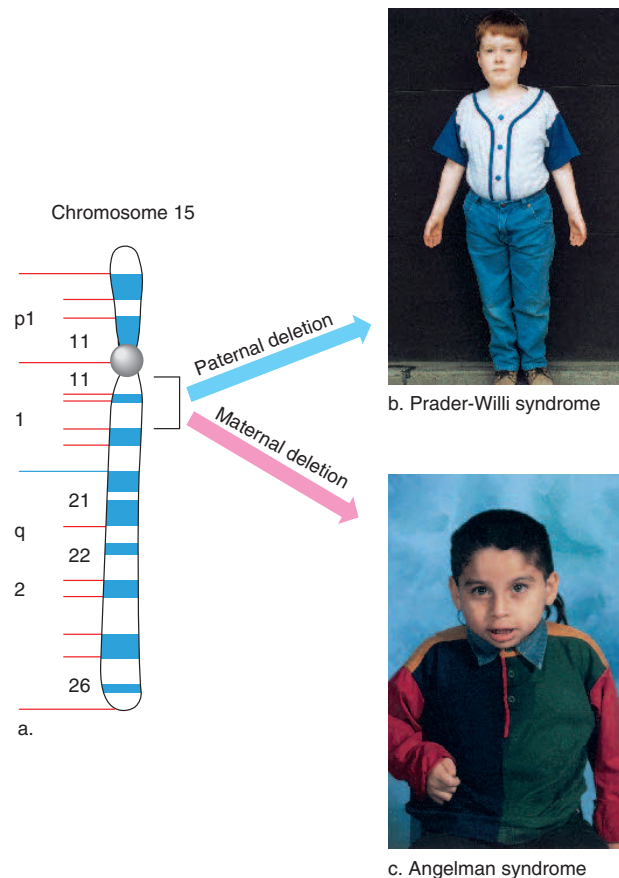


Figure 6.16 Prader-Willi and Angelman syndromes result when the non-imprinted copy of a gene is deleted.

(a) Two distinct syndromes result from missing genetic material in the same region of chromosome 15. (b) Tyler has Prader-Willi syndrome, due to a deletion in the copy of the chromosome he inherited from his father. Note his small hands. (c) This child has Angelman syndrome, caused by a deletion in the chromosome 15 that he inherited from his mother. He is mentally retarded.

who inherits both copies of a gene from one parent and none from the other. (Chapter 13 discusses uniparental disomy further.)

Key Concepts

1. In genomic imprinting, the phenotype differs depending on whether a gene is inherited from the mother or the father.
2. Methyl groups may bind to DNA and temporarily suppress gene expression in a pattern determined by the individual's sex.
3. Imprinting may be a normal process in mammalian embryos.

Summary

6.1 Our Sexual Selves

1. Sexual identity includes sex chromosome makeup; gonadal specialization; phenotype (reproductive structures); and gender identity.
2. The human male is the **heterogametic sex**, with an X and a Y chromosome. The female, with two X chromosomes, is the **homogametic sex**.
3. The human Y chromosome includes two pseudoautosomal regions and a large, male-specific region that does not recombine. Y-linked genes may correspond to X-linked genes, be similar to them, or be unique. Palindromic DNA sequences or inverted repeats can promote gene loss on the Y.
4. If the *SRY* gene is expressed, undifferentiated gonads develop as testes. If *SRY* is not expressed, the gonads develop as ovaries, under the direction of other genes.
5. Starting about 8 weeks after fertilization, sustentacular cells in the testes secrete anti-Müllerian hormone, which prevents development of female structures, and interstitial cells produce testosterone, which triggers development of the epididymides, ductus deferentia, seminal vesicles, and ejaculatory ducts.
6. Testosterone converted to DHT controls development of the urethra, prostate gland, penis, and scrotum. If *SRY* is not turned on, the Müllerian ducts continue to develop into female reproductive structures.
7. Evidence points to an inherited component to homosexuality.
8. **Sex ratio** is the number of males divided by the number of females multiplied by 1,000, for people of a particular age. Sex ratios are skewed by interfering with pregnancy outcomes.

6.2 Traits Inherited on Sex Chromosomes

9. Y-linked traits are rare and are passed from fathers to sons only.

10. Males are **hemizygous** for genes on the X chromosome and express phenotypes associated with these genes because they do not have another allele on a homolog. An X-linked trait passes from mother to son because he inherits his X chromosome from his mother and his Y chromosome from his father.
11. An X-linked allele may be dominant or recessive. X-linked dominant traits are more devastating to males than to females.

6.3 Sex-Limited and Sex-Influenced Traits

12. **Sex-limited traits** may be autosomal or sex-linked, but they only affect one sex because of anatomical or hormonal gender differences.
13. A **sex-influenced gene** is dominant in one sex but recessive in the other.

6.4 X Inactivation

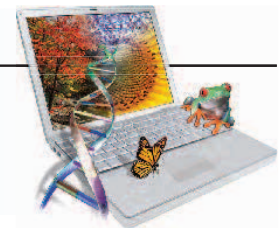
14. **X inactivation** shuts off one X chromosome in each cell in female mammals, making them mosaics for heterozygous genes on the X chromosome. This phenomenon evens out the dosages of genes on the sex chromosomes between the sexes.
15. A female who expresses the phenotype corresponding to an X-linked gene she carries is a **manifesting heterozygote**.

6.5 Genomic Imprinting

16. In **genomic imprinting**, the phenotype corresponding to a particular genotype differs depending on whether the parent who passes the gene is female or male.
17. Imprints are erased during meiosis and reassigned based on the sex of a new individual.
18. Methyl groups that temporarily suppress gene expression are the physical basis of genomic imprinting.

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Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

1. How is sex expressed at the chromosomal, gonadal, phenotypic, and gender identity levels?
2. How do genes in the pseudoautosomal region of the Y chromosome differ from genes in the male-specific region (MSY)?
3. What are the phenotypes of the following individuals?
 - a. a person with a mutation in the *SRY* gene, rendering it nonfunctional
 - b. a normal XX individual
 - c. an XY individual with a block in testosterone synthesis
4. List the events that must take place for a fetus to develop as a female.
5. Cite evidence that may point to a hereditary component to homosexuality.

- Why is it unlikely one would see a woman who is homozygous for an X-linked dominant condition?
- What is the basis of sex ratio at birth?
- Traits that appear more frequently in one sex than the other may be caused by genes that are inherited in an X-linked, sex-limited, or sex-influenced fashion. How might you distinguish among these possibilities in a given individual?
- Why are male calico cats very rare?
- How might X inactivation cause patchy hairiness in women who have congenital generalized hypertrichosis, even though the disease-causing allele is dominant?
- How does X inactivation even out the “doses” of X-linked genes between the sexes?
- Cite evidence that genetic contributions from both parents are necessary for normal prenatal development.
- Prader-Willi and Angelman syndromes are more common in children conceived with certain assisted reproductive technologies (*in vitro* fertilization and intracytoplasmic sperm injection) than among the general population. What process may these procedures disrupt?

Applied Questions

- To answer the following questions, consider these population data on sex ratios:

Selected sex ratios at birth		Selected sex ratios after age 65	
Nation	Sex ratio	Nation	Sex ratio
Costa Rica	970	Rwanda	620
Tanzania	1,000	South Africa	630
Liechtenstein	1,010	France	700
South Africa	1,020	United States	720
United States	1,050	Qatar	990
Sweden	1,060	Montserrat	1,060
Italy	1,070	Bangladesh	1,160
China	1,130	Nigeria	990

- In Rwanda, South Africa, France, and the U.S., males die, on average, significantly younger than females. What types of information might explain the difference?
- In Costa Rica, how many males at birth are there for every 100 females?
- In which country listed do males tend to live the longest?

- In severe Hunter syndrome, lack of the enzyme iduronate sulfate sulfatase leads to buildup of certain carbohydrates swelling the liver, spleen, and heart. In mild cases, deafness may be the only symptom. Intellect is usually unimpaired, and life span can be normal. Hunter syndrome is X-linked recessive. A man with mild Hunter syndrome has a child with a woman who is a carrier.
 - What is the probability that a son inherits Hunter syndrome?
 - What is the chance that a daughter inherits Hunter syndrome?
 - What is the chance that a daughter is a carrier?
- Amelogenesis imperfecta (MIM 301200) is X-linked dominant. Affected males have extremely thin enamel on each tooth. Female carriers have grooved teeth from uneven deposition of enamel. Why might the phenotype differ between the sexes?

Web Activities

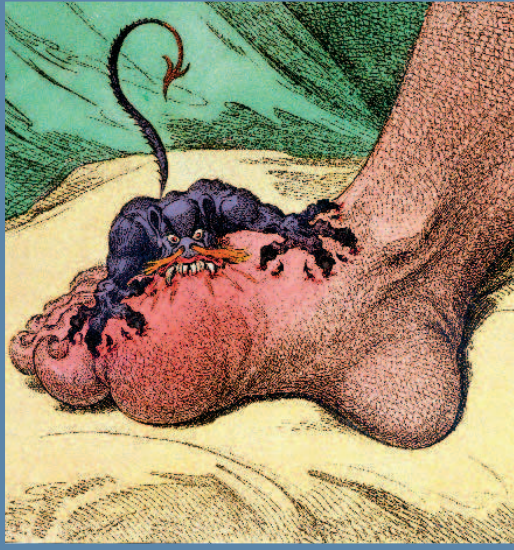
- Visit the National Center for Biotechnology Information (NCBI) website. Identify an X-linked disorder, then find it in MIM and describe it.
- At the Imprinted Gene Catalogue website, click on “search by species name” and then click on “complete list.” Find two

disorders that involve imprinting, one transmitted from the mother and one from the father, and use MIM to describe them.

Case Studies and Research Results

- For each case description, identify the principle at work from the list that follows. More than one answer per case may apply.
 - Y-linked inheritance
 - X-linked recessive
 - X-linked dominant inheritance
 - Sex-limited inheritance
 - Sex-influenced inheritance
 - X inactivation or manifesting heterozygote
 - Uniparental disomy
 - Imprinting abnormality
 - In a three-generation family, sixteen members have speech-language disorder (MIM 602081) and cannot speak. The gene that is mutant is called *FOXP2*. The speechless family members inherited both copies of the gene from their mothers and none from their fathers.

- b. Six-year-old LeQuan inherited Fabry disease (MIM 301500) from his mother, who is a heterozygote for the causative mutation. The gene, on the X chromosome, encodes a lysosomal enzyme. LeQuan would die before age 50 of heart failure, kidney failure, or a stroke, but fortunately he can be treated with twice-monthly infusions of the enzyme. His mother, Echinecea, recently began experiencing recurrent fevers, a burning pain in her hands and feet, a rash, and sensitivity to cold. She is experiencing mild Fabry disease.
- c. The Chandler family has many male members who have a form of retinitis pigmentosa (RP) in which the cells that capture light energy in the retina degenerate, causing gradual visual loss. Several female members of the family presumed to be carriers because they have affected sons are tested for RP genes on chromosomes 1, 3, 6, and the X, but do not carry these RP genes. Many years ago, Rachel married her cousin Ross, who has the family's form of RP. They had six children. The three sons are all affected, but their daughters all have normal vision.
- d. Simon's mother and her sister are breast cancer survivors, and their mother died of the disease. Simon's sister Maureen has a genetic test and learns that she, too, has inherited the *BRCA1* gene. Simon has two daughters, but doesn't want to be tested because he thinks a man cannot transmit a trait that affects a body part that is more developed in females.
- e. Tribbles are extraterrestrial mammals that long ago invaded a starship on the television program *Star Trek*. A gene called *frizzled* causes kinky hair in female tribbles who inherit just one allele. However, two mutant alleles must be inherited for a male tribble to have kinky hair.
- f. Prozac died at age 16 of Lowe syndrome (MIM 309000). He was slightly mentally retarded, had visual problems (cataracts and glaucoma), seizures, poor muscle tone, and progressive kidney failure, which was ultimately fatal. His sister Lunesta is pregnant, and wonders whether she is a carrier of the disease that killed her brother. She remembers a doctor saying that her mother Yaz was a carrier. Lunesta's physician determines that she is a carrier because she has cataracts, which is a clouding of the lenses. It has not yet affected her vision. When a prenatal test reveals that Lunesta's fetus is a female, her doctor tells her not to worry about Lowe syndrome.
- g. Mating among Texas field crickets depends upon females responding to a male mating call. The sounds must arrive at a particular frequency to excite the females, who do not sing back in response. However, females can pass on a trait that confers frequency of singing.
- h. When Winthrop was a baby, he was diagnosed with "failure to thrive." At 14 months of age, he suddenly took an interest in food, and his parents couldn't feed him fast enough. By age 4, Winthrop was obese, with disturbing behavior. He was so hungry that after he'd eaten his meal and everyone else's leftovers, he'd hunt through the garbage for more. Finally a psychiatrist who had a background in genetics diagnosed Prader-Willi syndrome. Testing showed that the allele for the Prader-Willi gene that Winthrop had inherited from his father was abnormally methylated.
- i. Certain breeds of dogs have cryptorchidism, in which the testicles do not descend into the scrotum. The trait is passed through females.
7. Reginald has mild hemophilia A that he can control by taking a clotting factor. He marries Lydia, whom he met at the hospital where he and Lydia's brother, Marvin, receive their treatment. Lydia and Marvin's mother and father, Emma and Clyde, do not have hemophilia. What is the probability that Reginald and Lydia's son will inherit hemophilia A?
8. Harold works in a fish market, but the odor does not bother him because he has anosmia (MIM 301700), an X-linked recessive lack of sense of smell. Harold's wife, Shirley, has a normal sense of smell. Harold's sister, Maude, also has a normal sense of smell, as does her husband, Phil, and daughter, Marsha, but their identical twin boys, Alvin and Simon, cannot detect odors. Harold and Maude's parents, Edgar and Florence, can smell normally. Draw a pedigree for this family, indicating people who must be carriers of the anosmia gene.



Gout was once associated with royalty because only they could afford the kidney pies, liver, brains, and scallops that cause uric acid to be deposited in the joints, mostly those of the big toe. The painful condition can develop if a genetically susceptible individual indulges in these foods.

CHAPTER

7

Multifactorial Traits

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- 7.2 Polygenic Traits Are Continuously Varying**
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 - Empiric Risk
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- 7.4 Genome-Wide Association Studies**
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Hold the Anchovies: Genes and Diet Cause Gout

The pain of gout in one's big toe is sudden and excruciating. Two to six million people in the United States have the condition, which results from deposition of uric acid crystals in a joint. Once known as "the disease of kings" because it was seen mostly among royalty, we now know that gout occurs when a genetically susceptible individual eats a diet rich in organ meats and fish. These foods are mostly muscle tissue, packed with mitochondria that have abundant nucleic acid purine bases that form uric acid.

The Egyptians first described gout in 2640 B.C., and in the fifth century B.C., Hippocrates called it "the unwalkable disease." King Charles I of Spain gave up his vast empire in 1556 due to the painful condition. In 2006, Spanish researchers confirmed the king's diagnosis by detecting uric acid deposits in the end joint of a finger that, for reasons unknown, had been preserved in a small box apart from the rest of the king.

Today gout provides a great example of the power of genome-wide association studies to identify inherited susceptibility to a painful condition. Such a study seeks associations by mining very large samples. To search for gout susceptibility alleles, researchers scanned the genomes of 26,714 individuals enrolled in three major health studies in the United States and the Netherlands for 500,000 SNPs (single-base sites in the genome that vary in at least 1 percent of a population). Just three of the half million SNPs were common to those individuals who had high uric acid levels in the blood, which indicates a tendency toward developing gout. The SNPs were very rare in others.

The implicated SNPs are in or near genes that make sense—all three genes affect the ability of kidney tubules to excrete uric acid. Each person has two copies of each of the three susceptibility genes, represented by six SNPs. For a person with none of the telltale SNPs, the risk of developing gout is 1 to 2 percent, but for someone with all six, the risk is 8 to 18 percent. The next step is to develop a susceptibility test based on these gene variants to identify people who should not feast on chopped liver and herring.

7.1 Genes and the Environment Mold Traits

A woman who is a prolific writer has a daughter who becomes a successful novelist. An overweight man and woman have obese children. A man whose father suffers from alcoholism has the same problem. Are these characteristics—writing talent, obesity, and alcoholism—inherited or learned? These traits, and nearly all others, are not the result of an “either/or” mechanism, but reflect the input of many genes as well as environmental influences. Even single-gene disorders are modified by environmental factors and/or other genes. A child with cystic

fibrosis, for example, has inherited a single-gene disorder, but her experiences reflect which variants of the gene she has, other genes that affect her immune system, the pathogens to which she is exposed, and the quality of the air she breathes. Some geneticists use the term “oligogenic disorder” for a condition caused by one major gene and smaller effects of other genes. This chapter looks beyond the rare, single-gene characteristics and disorders of the past several chapters to the more common characteristics that represent input from many genes, and the tools used to study them.

A trait can be described as either single-gene (or Mendelian or monogenic) or **polygenic**. As its name implies, a polygenic trait reflects the activities of more than one gene. **Figure 7.1** compares single-gene traits with those that result from the actions of multiple genes. Both single-gene and polygenic traits can also be **multifactorial**, which means they are influenced by the environment. Pure polygenic traits—those not influenced by the environment—are very rare. Eye color, discussed in chapter 4, is close to being purely polygenic.

Polygenic multifactorial traits include common ones, such as height, skin color, body weight, many illnesses, and behavioral conditions and tendencies. Behavioral traits are not inherently different from other types of traits; they involve the functioning of the brain, rather than another organ. Chapter 8 discusses them. A more popular term for “multifactorial” is

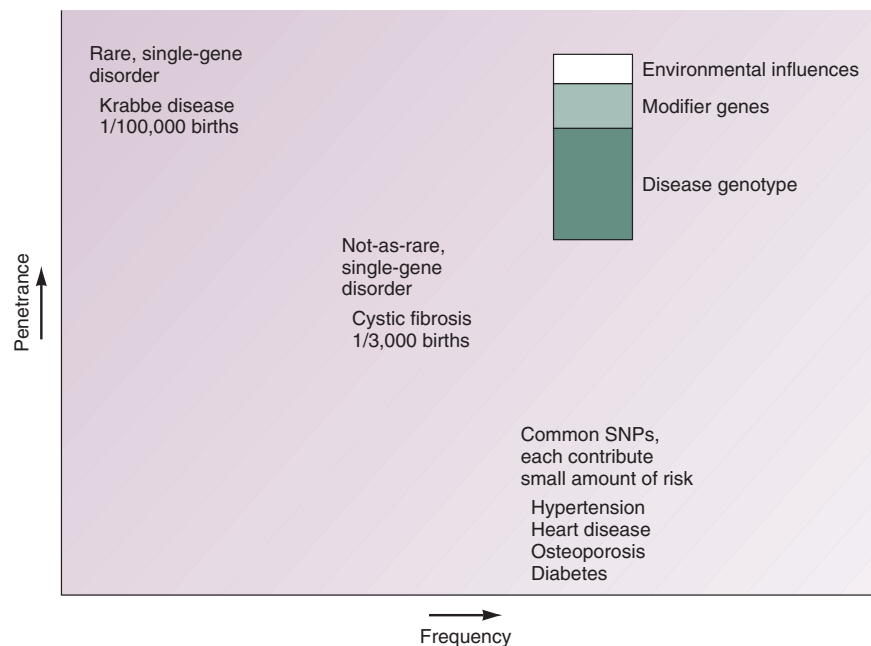


Figure 7.1 Anatomy of a trait. Few, if any, genes act alone. Genes influence each other and respond to environmental stimuli by their levels of expression. We classify genetic disorders by their penetrance (percent of individuals with a genotype who have the associated phenotype) and frequency (how common they are). Mutations and SNPs are both changes in DNA, but mutations are rare and often associated with single-gene disorders. SNPs are more common, and many may contribute in a small way to a trait or condition. This figure compares an extremely rare single-gene disease, a more common single-gene disease, and SNPs that are associated with increased disease risk, in terms of penetrance and frequency. The inset depicts a hypothetical single-gene disorder and the contribution of modifier genes and the environment to the phenotype.

complex, but we use multifactorial here because it is more precise and is not confused with the general definition of “complex.” The genes of a multifactorial trait are not more complicated than others. They follow Mendel’s laws, but expression of any one gene is more difficult to predict because of the combined actions of genes and the environment.

Lung cancer caused by smoking illustrates the complexity of multifactorial traits. Variants of genes that increase the risk of becoming addicted to nicotine and of developing cancer come into play—but may not ever be expressed if a person never smokes or breathes polluted air.

A polygenic multifactorial condition reflects additive contributions of several genes. Each gene confers a degree of susceptibility, but the input of these genes is not necessarily equal. For example, three genes contribute significantly to the risk of developing type 2 diabetes mellitus, but other genes may exert smaller effects.

Different genes may contribute different aspects of a phenotype that was once thought to be due to the actions of a single gene. Consider migraine, a condition which many a sufferer will attest is more than just a headache. Studies have found that a gene on chromosome 1 contributes sensitivity to sound; a gene on chromosome 5 produces the pulsating headache and sensitivity to light; and a gene on chromosome 8 is associated with nausea and vomiting. In addition, certain environmental influences are well known to trigger migraine in some people. **Reading 7.1** takes a closer look at heart health, which reflects several multifactorial traits.

7.2 Polygenic Traits Are Continuously Varying

For a polygenic trait, the combined action of many genes often produces a “shades of grey” or “continuously varying” phenotype, also called a quantitative trait. DNA sequences that contribute to polygenic traits are called **quantitative trait loci**, or QTLs. A multifactorial trait is continuously varying if it is also polygenic. That is, it is the multi-gene component of the trait that contributes the continuing variation of the phenotype. The individual genes that confer a polygenic trait follow Mendel’s laws, but together they do not produce single-gene phenotypic ratios. They all contribute to the phenotype, but without being dominant or recessive to each other. Single-gene traits are instead discrete or qualitative, often providing an “all-or-none” phenotype such as “normal” versus “affected.”

A polygenic trait varies in populations, as our many nuances of hair color, body weight, and cholesterol levels demonstrate. Some genes contribute more to a polygenic trait than others. Within genes, alleles can have differing impacts depending upon exactly how they alter an encoded protein and how common they are in a population. For example, a mutation in the gene that encodes the receptor that takes LDL cholesterol into cells greatly raises blood serum cholesterol level. But because fewer than 1 percent of the individuals in most

populations have this mutation, it contributes very little to the variation in cholesterol level at the population level.

Although the expression of a polygenic trait is continuous, we can categorize individuals into classes and calculate the frequencies of the classes. When we do this and plot the frequency for each phenotype class, a bell-shaped curve results. Even when different numbers of genes affect the trait, the curve takes the same shape, as is evident in the following examples.

Fingerprint Patterns

The skin on the fingertips is folded into patterns of raised skin called dermal ridges that align to form loops, whorls, and arches. This pattern is a fingerprint. A technique called dermatoglyphics (“skin writing”) compares the number of ridges that comprise these patterns to identify and distinguish individuals (**figure 7.2**). Dermatoglyphics is part of genetics because certain disorders (such as Down syndrome) include unusual ridge patterns. Forensic fingerprint analysis is also an application of dermatoglyphics.

The number of ridges in a fingerprint is largely determined by genes, but also arises from the environment. During weeks 6 through 13 of prenatal development, the ridge pattern can be altered as the fetus touches the finger and toe pads to the wall of the amniotic sac. This early environmental effect explains why the fingerprints of identical twins, who share all genes, are in some cases not exactly alike.

We can quantify a fingerprint with a measurement called a total ridge count, which tallies the numbers of ridges in whorls, loops, or arches. The average total ridge count in a male is 145, and in a female, 126. Plotting total ridge count reveals the bell curve of a continuously varying trait.

Height

The effect of the environment on height is obvious—people who do not eat enough do not reach their genetic potential for height. Students lined up according to height, but raised in two different decades and under different circumstances, vividly reveal the effects of genes and the environment on this continuously varying trait. Part *a* of **figure 7.3** depicts students from 1920, and part *b*, students from 1997. Also note that the tallest people in the old photograph are 5’9”, whereas the tallest people in the more recent photograph are 6’5”. The difference is attributed to improved diet and better overall health.

Genome-wide association studies have identified dozens of genes that affect height. Such a study compares genetic markers (see table 5.3) in two large groups of individuals who, ideally, differ only in the characteristic of interest. An association between the trait and the marker is then inferred, which, after much more investigation, may uncover a gene that contributes to the characteristic. For example, a long-term study called the Northern Finland Birth Cohort is following various characteristics in 4,763 people born in 1966, recording their height 20 times so far. Certain patterns of SNPs in particular genes are found significantly more in individuals who experienced periods of rapid height increase. These findings may lead to insights about

Reading 7.1

Heart Health is Multifactorial

Many types of cells and processes must interact for the heart and vessels to effectively and continuously circulate blood. Therefore, it isn't surprising that many genes take part in the system's maintenance. Effects of the environment are great, too. Even single-gene cardiovascular diseases are affected by outside influences. For example, intake of vitamin K, necessary for blood to clot, influences the severity of single-gene clotting disorders.

Genes control cardiovascular functioning in several ways: transporting lipids; blood clotting; blood pressure; and how well cellular adhesion molecules enable white blood cells to stick to the walls of blood vessels. Lipids can only move in the circulation when bound to proteins to form large molecules called lipoproteins. Several genes encode the protein parts of lipoproteins, which are called apolipoproteins. Some types of lipoproteins carry lipids in the blood to tissues, where they are utilized, and other types of lipoproteins take lipids to the liver, where they are dismantled into biochemicals that the body can excrete more easily. One allele of a gene that encodes apolipoprotein E, called E4, increases the risk of a heart attack threefold in people who smoke. This is clear evidence that genes and environmental factors can interact in ways that cause illness.

Maintaining a healthy cardiovascular system requires a lipid balance between sufficient lipids inside cells but not an excess outside cells. Several dozen genes control lipid levels in the blood and tissues by specifying enzymes that process lipids, proteins that transport them, or receptor proteins that admit lipids into cells.

An enzyme, lipoprotein lipase, is particularly important in lipid metabolism. It lines the walls of the smallest blood vessels, where it breaks down fat packets released from the small intestine and liver. Lipoprotein lipase is activated by high-density lipoproteins (HDLs), and it breaks down low-density lipoproteins (LDLs). High HDL levels and low LDL levels are associated with a healthy cardiovascular system. Lipoprotein lipase also regulates fat cell size; fat cells contribute to obesity by enlarging, rather than dividing.

The fluidity of the blood is also critical to health. Overly active clotting factors or extra sticky white blood cells can induce formation of clots that block blood flow, usually in blood vessels in the heart or in the legs. Poor clotting causes dangerous bleeding. Because clotting factors are proteins, clotting is genetically controlled.

Although it is common to measure serum cholesterol level as a biomarker of cardiovascular disease, testing for C reactive protein may be more useful. This protein, which is in blood plasma, is part of the inflammatory response. High levels of it are associated with increased risk of heart attack due to inflammation in the arteries. However, elevation of C reactive protein could have other causes.

Genetic test panels detect multiple alleles in dozens of genes that cause or contribute to cardiovascular disease. More than 50 genes regulate blood pressure. In addition, DNA microarrays can monitor gene expression, assessing many contributing factors. For example, one gene expression microarray test can indicate which cholesterol-lowering drugs are most likely to be effective and without side effects for a particular individual. The premise behind the value of such information is that people have composite genetic risks based on the small contributions of several genes—the essence of polygenic inheritance.

Computer analysis of multigene tests accounts for environmental factors (**table 1**). Some risk factors are controllable with lifestyle changes, such as exercising, not smoking, and maintaining a healthy weight. **Figure 1** shows an artery blocked by fatty plaque. Diet and medication can counter an inherited tendency to deposit cholesterol-rich material on the interior linings of arteries.

Table 1

Risk Factors for Cardiovascular Disease

Uncontrollable	Controllable
Age	Fatty diet
Male sex	Hypertension
Genes	Smoking
Lipid metabolism	High serum cholesterol
Apolipoproteins	Low serum HDL
Lipoprotein lipase	High serum LDL
Blood clotting	Stress
Fibrinogen	Insufficient exercise
Clotting factors	Obesity
Inflammation	
C reactive protein	
Homocysteine metabolism	Diabetes
Leukocyte adhesion	

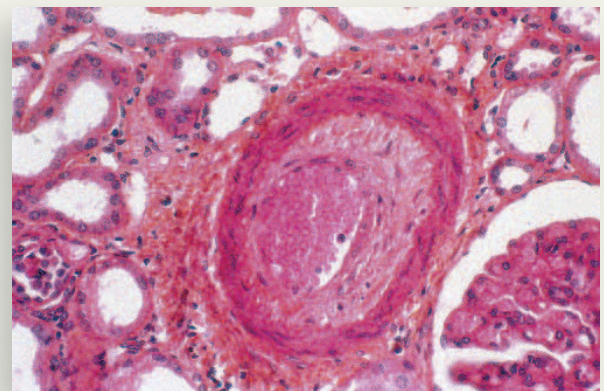


Figure 1 Cardiovascular disease. Genetic and dietary factors contribute to clogged arteries.

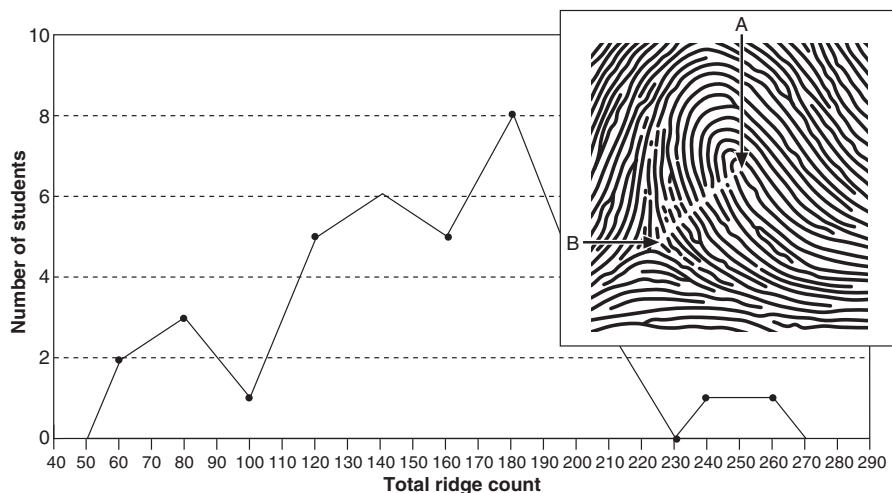


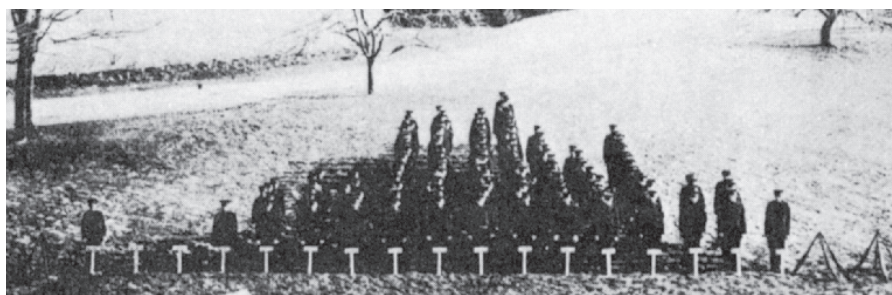
Figure 7.2 Anatomy of a fingerprint. Total ridge counts for a number of individuals, plotted on a bar graph, form an approximate bell-shaped curve. The number of ridges between landmark points A and B on this loop pattern is 12. Total ridge count includes the number of ridges on all fingers.

which genes affect height at different ages. Section 7.4 discusses these types of studies in more detail.

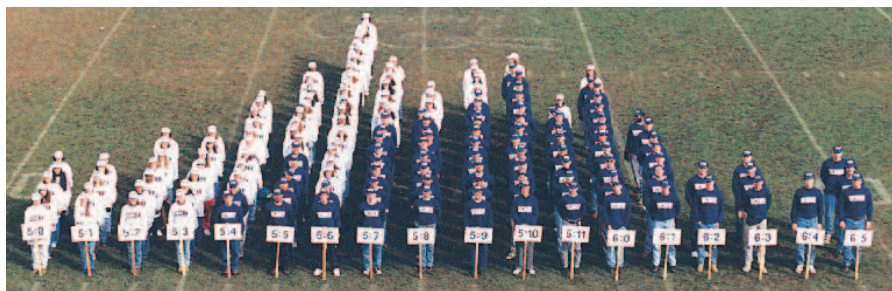
Skin Color

More than 100 genes affect pigmentation in skin, hair, and the irises. Melanin pigments color the skin to different degrees in different individuals. In the skin, as in the iris (see chapter 4),

melanocytes contain melanin in melanosomes. Melanocytes extend between the tile-like skin cells, distributing pigment granules through the skin layers. Some melanin exits the melanocytes and enters the hardened cells in the skin's upper layers. Here the melanin breaks into pieces, and as the skin cells are pushed up toward the skin's surface, the melanin bits provide color. The pigment protects against DNA damage from ultraviolet radiation, and exposure to the sun increases melanin synthesis.



a.



b.

Figure 7.3 The inheritance of height. The photograph in (a) illustrates the continuously varying nature of height. In the photo, taken around 1920, 175 cadets at the Connecticut Agricultural College lined up by height. (b) In 1997, professor Linda Strausbaugh asked her genetics students at the school (today the University of Connecticut at Storrs) to re-create the scene.

Figure 7.4 shows a three-gene model for human skin color—an oversimplification of this highly variable trait that does, however, illustrate how many genes can contribute to this trait.

Although people come in a wide variety of hues, we all have about the same number of melanocytes per unit area of skin. However, people differ in melanosome number, size, and density of distribution. Differences in skin color arise from the number and distribution of melanin pieces in the skin cells in the uppermost layers.

Skin color is one trait used to distinguish race. However, the definition of race based largely on skin color is more a social construct than a biological concept, for skin color is but one of thousands of traits whose frequencies vary in different populations. From a genetic perspective, when referring to other types of organisms, races are groups within species that are distinguished by different allele frequencies. Humans are actually a lot less variable in appearance than other mammals, although it may seem to us that all chimps look alike. Although we tend to classify people by skin color because it is an obvious visible way to distinguish individuals, skin color is *not* a reliable indicator of ancestry.

When many genes are considered, two people with very dark skin may be less alike than either is to another person with very light skin. For example, sub-Saharan Africans and Australian aborigines have dark skin, but are very dissimilar in other inherited characteristics. Their dark skins may reflect adaptation to life in a sunny, tropical climate rather than recent shared ancestry. Overall, 93 percent of varying inherited traits are no more common in people of one skin color than any other.

Testing DNA indicates that biologically speaking, it makes more sense to classify people by ancestry rather than by the color of their skin. In one telling investigation, 100 students in a sociology class in “Race and Ethnic Relations” at Pennsylvania State University had their DNA tested for percent contribution from “European white,” “black African,” “Asian,” and “Native American” gene variants that are more common in these groups. No student was pure anything, and many were quite surprised at what their DNA revealed about their ancestry. One student, a light-skinned black, learned that genetically he is 52 percent black African and 48 percent European white:

approximately half black, half white. Another student who considered herself black was actually 58 percent white European. The U.S. census, in recognition of the complexity of classifying people into races based on skin color, began to allow “mixed race” as a category in 2000. Many of us fall into this category, including President Obama.

Although in a genetic sense the concept of race based on skin color has little meaning, in a practical sense, racial groups do have different incidences of certain diseases. This reflects the tendency to choose partners within a group, which retains certain alleles. However, racial differences in disease prevalence may also result from social inequities, such as some groups’ limited access to good nutrition or health care. Observations that populations of particular races have a higher incidence of certain illnesses have fueled “race-based prescribing.” For example, certain hypertension and heart disease drugs are specifically marketed to African Americans, who have a higher incidence of these conditions than do people in other groups. **Table 7.1** lists some drugs that seem to be more effective among either Americans of European descent or Americans of African descent.

Offering medical treatments based on skin color may make sense on a population level, but on the individual level it may lead to errors. A white person might be denied a drug that would work, or a black person given one that doesn’t, if the treatment decision is based on a superficial trait not directly related to how the body responds to a particular drug.

Prescribing drugs is beginning to be based on personal genotypes that determine whether or not a particular drug will work or have side effects, rather than by the color of a person’s skin. For example, a gene called *MDR* (for multi-drug resistance) encodes a protein that pumps poisons out of certain white blood cells and intestinal lining cells. When a gene variant results in a pump that works too well, the protein recognizes drugs used to treat cancer, AIDS, and other conditions as toxins, sending them out of the cell. Researchers have found this protein variant in 83 percent of West Africans, 61 percent of African Americans, 26 percent of Caucasians, and 34 percent of Japanese. That is, a person in whom these drugs do not work could come from any of these populations.

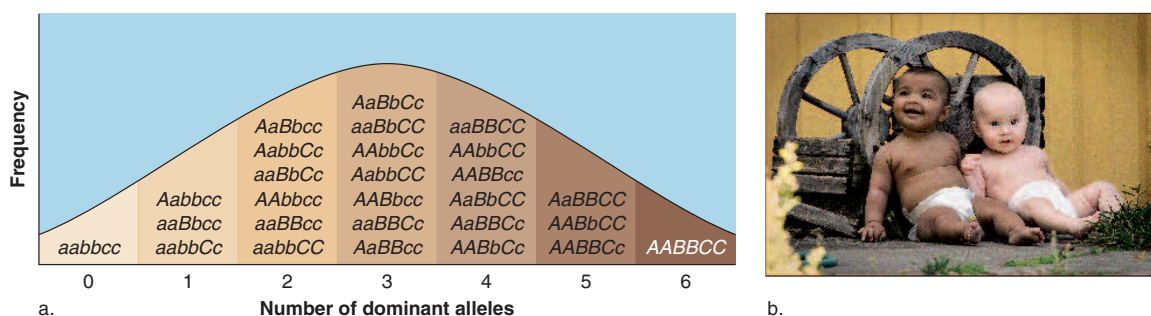


Figure 7.4 Variations in skin color. (a) A model of three genes, with two alleles each, can explain broad hues of human skin. In actuality, this trait likely involves many more than three genes. (b) Humans come in a great variety of skin colors. Skin color genes can sort themselves out in interesting ways. These beautiful young ladies, Alicia and Jasmin, are twins! Their father is German and their mother is Jamaican-English.

Table 7.1 Different Drug Responses Among European Americans and African Americans

Researchers have compared the effects of many drugs in different racial or ethnic groups. Most drugs tested either show no difference, or do show a difference but the physiological basis isn't known. For some drugs, one group will tend to respond to lower doses or in greater numbers than another. Some of these drugs are listed here. The difference in response is associated with inheriting particular gene variants or SNPs. (EA stands for European American and AA stands for African American.)

Drug Class/Name	Disorder	More Effective In:	
		EA	AA
ACE inhibitor/Enalapril	hypertension	✓	
Antipsychotic/Clozapine	psychosis		✓
Antiviral/Alpha interferon	hepatitis	✓	
Beta blocker/Propranolol	hypertension	✓	
Calcium channel blocker/Diltiazem	hypertension		✓
Insulin	diabetes mellitus	✓	
Thiazide diuretic	hypertension		✓
Vasodilator combination/BiDil	congestive heart failure		✓

MDR genotype, then, can be used to prescribe certain drugs only for individuals whose cells would not pump the drugs out. MDR genotype is a more biologically meaningful basis for prescribing a drug than skin color.

In an even more compelling study, researchers cataloged 23 markers for genes that control drug metabolism in 354 people representing eight races: black (Bantu, Ethiopian, and Afro-Caribbean,), white (Norwegian, Armenian, and Ashkenazi Jews), and Asian (Chinese and New Guinean). The genetic markers fell into four very distinct groups that predict which of several blood thinners, chemotherapies, and painkillers will be effective—and these response groups did not at all match the traditional racial groups.

The premises behind race-based prescribing are far more complex than black versus white. Although some genes and their variants are not distributed along racial lines, such as the 23 markers of drug metabolism just discussed, others apparently are. This is the case for a gene that encodes an enzyme called leukotriene A4 hydrolase. The enzyme is necessary to produce leukotrienes, which inflame arteries as part of the immune response to infection. Excess leukotrienes increase the risk of heart attack. An allele present in European Americans and European populations for many years increases heart attack risk only slightly. Researchers hypothesize that enough time has passed that variants of other genes that temper the negative effects of excess leukotrienes have accumulated in the genomes of these light-skinned groups. The overactive leukotriene A4 hydrolase allele, however, has only recently been introduced into the African American population. Without enough time for genetic protection to have arisen, the excess leukotrienes

elevate risk of heart attack among African Americans five-fold compared to 16 percent elevation among whites.

Key Concepts

1. Polygenic traits are determined by more than one gene and vary continuously in expression.
2. Multifactorial traits are determined by a combination of a gene or genes and the environment.
3. A bell curve describes the distribution of phenotypic classes of a polygenic trait.

7.3 Traditional Approaches To Investigating Multifactorial Traits

Predicting recurrence risks for polygenic traits is much more challenging than doing so for single-gene traits. This section reviews traditional approaches to evaluating polygenic multifactorial traits, and the next section examines genome-wide association studies.

Empiric Risk

Using Mendel's laws, it is possible to predict the risk that a single-gene trait will recur in a family from knowing the mode of inheritance—such as autosomal dominant or recessive. To predict the chance that a polygenic multifactorial trait will occur

in a particular individual, geneticists use **empiric risk**, which is based on incidence in a specific population. **Incidence** is the rate at which a certain event occurs, such as the number of new cases of a particular disorder diagnosed per year in a population of known size. **Prevalence** is the proportion or number of individuals in a population who have a particular disorder at a specific time, such as during one year.

Empiric risk is not a calculation, but a population statistic based on observation. The population might be broad, such as an ethnic group or community, or genetically more well-defined, such as families that have a particular disease. Empiric risk increases with the severity of the disorder, the number of affected family members, and how closely related a person is to affected individuals. As an example, consider using empiric risk to predict the likelihood of a child being born with a neural tube defect (NTD). In the United States, the overall population risk of carrying a fetus with an NTD is about 1 in 1,000 (0.1 percent). For people of English, Irish, or Scottish ancestry, the risk is about 3 in 1,000. However, if a sibling has an NTD, no matter what the ethnic group, the risk of recurrence increases to 3 percent, and if two siblings are affected, the risk to a third child is even greater. By determining whether a fetus has any siblings with NTDs, a genetic counselor can predict the risk to that fetus, using the known empiric risk.

If a trait has an inherited component, then it makes sense that the closer the relationship between two individuals, one of whom has the trait, the greater the probability that the second individual has the trait, too, because they have more genes in common. Studies of empiric risk support this logic. **Table 7.2** summarizes empiric risks for relatives of individuals with cleft lip (**figure 7.5**).

Because empiric risk is based solely on observation, we can use it to derive risks for disorders with poorly understood transmission patterns. For example, certain multifactorial disorders affect one sex more often than the other. Pyloric stenosis, an overgrowth of muscle at the juncture between the stomach and the small intestine, is five times more common among males than females. The condition must be corrected surgically shortly after birth, or the newborn will be unable to digest foods. Empiric data show that the risk of recurrence for



Figure 7.5 Cleft lip. Cleft lip is more likely in a person who has a relative with the condition. This child has had corrective surgery.

the brother of an affected brother is 3.8 percent, but the risk for the brother of an affected sister is 9.2 percent. An empiric risk, then, is based on real-world observations—the mechanism of the illness or its cause need not be known.

Table 7.2 Empiric Risk of Recurrence for Cleft Lip	
Relationship to Affected Person	Empiric Risk of Recurrence
Identical twin	40.0%
Sibling	4.1%
Child	3.5%
Niece/nephew	0.8%
First cousin	0.3%
General population risk (no affected relatives)	0.1%

Heritability

As Charles Darwin noted, some of the variation of a trait is due to inborn differences in populations, and some to differences in environmental influences. A measurement called **heritability**, designated *H*, estimates the proportion of the phenotypic variation for a particular trait that is due to genetic differences in a certain population at a certain time. The distinction between empiric risk and heritability is that empiric risk could result from nongenetic influences, whereas heritability focuses on the genetic component of the variation in a trait.

Figure 7.6 outlines the factors that contribute to observed variation in a trait. Heritability equals 1.0 for a trait whose variability is completely the result of gene action, such as in

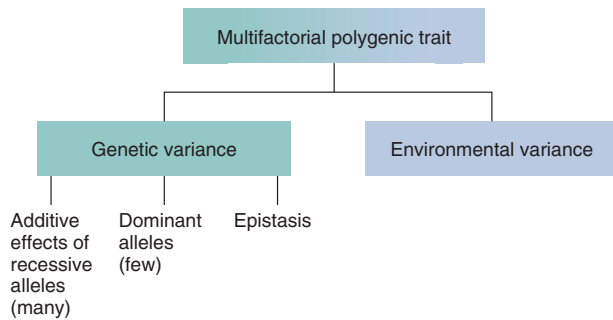


Figure 7.6 Heritability estimates the genetic contribution to the variability of a trait. Observed variance in a polygenic, multifactorial trait or illness reflects genetic and environmental contributions.

a population of laboratory mice whose environment is controlled. Without environmental variability, genetic differences determine expression of the trait in the population. Variability of most traits, however, reflects a combination of differences among genes and environmental components. **Table 7.3** lists some traits and their heritabilities.

Heritability changes as the environment changes. For example, the heritability of skin color would be higher in the winter months, when sun exposure is less likely to increase melanin synthesis. The same trait may be highly heritable in two populations, but certain variants much more common in one group due to long-term environmental differences. Populations in equatorial Africa, for example, have darker skin than sun-deprived Scandinavians.

Researchers use several statistical methods to estimate heritability. One way is to compare the actual proportion of pairs of people related in a certain manner who share a particular trait, to the expected proportion of pairs that would share it if it were inherited in a Mendelian fashion. The expected

Trait	Heritability
Clubfoot	0.8
Height	0.8
Blood pressure	0.6
Body mass index	0.5
Verbal aptitude	0.7
Mathematical aptitude	0.3
Spelling aptitude	0.5
Total fingerprint ridge count	0.9
Intelligence	0.5–0.8
Total serum cholesterol	0.6

Relationship	Degree of Relationship	Percent Shared Genes (Coefficient of Relatedness)
Sibling to sibling	1°	50% (1/2)
Parent to child	1°	50% (1/2)
Uncle/aunt to niece/nephew	2°	25% (1/4)
Grandparent to grandchild	2°	25% (1/4)
First cousin to first cousin	3°	12 1/2% (1/8)

proportion is derived by knowing the blood relationships of the individuals and using a measurement called the **coefficient of relatedness**, which is the proportion of genes that two people related in a certain way share (**table 7.4**).

A parent and child share 50 percent of their genes, because of the mechanism of meiosis. Siblings share on average 50 percent of their genes, because they have a 50 percent chance of inheriting each allele for a gene from each parent. Genetic counselors use the designations of primary (1°), secondary (2°), and tertiary (3°) relatives when calculating risks (**table 7.4** and **figure 7.7**). For extended or complicated pedigrees, the value of 1 in 2 or 50 percent between siblings and between parent-child pairs can be used to trace and calculate the percentage of genes shared between people related in other ways.

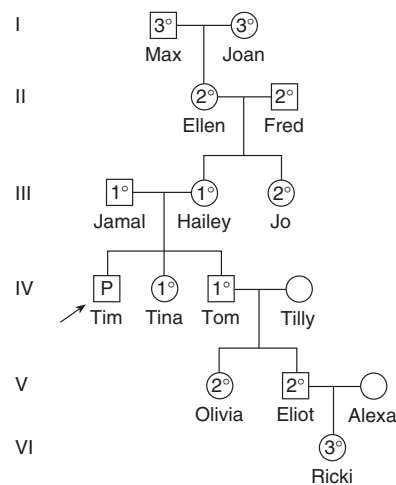


Figure 7.7 Tracing relatives. Tim has an inherited illness. A genetic counselor drew this pedigree to explain the approximate percentage of genes Tim shares with relatives. This information can be used to alert certain relatives to their risk.

("P" is the proband, or affected individual who initiated the study. See **table 7.4** for definitions of 1°, 2°, and 3° relationships.)

If the heritability of a trait is very high, then of a group of 100 sibling pairs, nearly 50 would be expected to have the same phenotype, because siblings share on average 50 percent of their genes. Height is a trait for which heritability reflects the environmental influence of nutrition. Of 100 sibling pairs in a population, for example, 40 might be the same number of inches tall. Heritability for height among this group of sibling pairs is .40/.50, or 80 percent, which is the observed phenotypic variation divided by the expected phenotypic variation if environment had no influence.

Genetic variance for a polygenic trait is mostly due to the additive effects of recessive alleles of different genes. For some traits, a few dominant alleles can greatly influence the phenotype, but because they are rare, they do not contribute greatly to heritability. This is the case for heart disease caused by a faulty LDL receptor. Epistasis (interaction between alleles of different genes) can also influence heritability. To account for the fact that different genes affect a phenotype to differing degrees, geneticists calculate a “narrow” heritability that considers only additive recessive effects, and a “broad” heritability that also considers the effects of rare dominant alleles and epistasis. For LDL cholesterol level, for example, the narrow heritability is 0.36, but the broad heritability is 0.96, reflecting the fact that a rare dominant allele has a large impact.

Understanding multifactorial inheritance is important in agriculture. A breeder needs to know whether genetic or environmental influences contribute to variability in such traits as birth weight, milk yield, and egg hatchability. It is also valuable to know whether the genetic influences are additive or epistatic. The breeder can control the environment by adjusting the conditions under which animals are raised and crops grown, and control genetic effects by setting up crosses between particular individuals.

Studying multifactorial traits in humans is difficult, because information must be obtained from many families. Two special types of people, however, can help geneticists to tease apart the genetic and environmental components of the variability of multifactorial traits—adopted individuals and twins.

Key Concepts

1. Empiric risk applies population incidence data to predict risk of recurrence for a multifactorial trait or disorder.
2. Heritability measures the genetic contribution to the variability of a multifactorial trait; it is specific to a particular population at a particular time.
3. Coefficient of relatedness, the proportion of genes that individuals related in a certain way are expected to share, is used to calculate heritability.

Adopted Individuals

A person adopted by people who are not blood relatives shares environmental influences, but typically not many genes, with the adoptive family. Conversely, adopted individuals share

genes, but not the exact environment, with their biological parents. Therefore, biologists assume that similarities between adopted people and adoptive parents reflect mostly environmental influences, whereas similarities between adoptees and their biological parents reflect mostly genetic influences. Information on both sets of parents can reveal how heredity and the environment contribute to a trait.

Many early adoption studies used a database of all adopted children in Denmark and their families from 1924 to 1947. One study examined correlations between causes of death among biological and adoptive parents and adopted children. If a biological parent died of infection before age 50, the adopted child was five times more likely to die of infection at a young age than a similar person in the general population. This may be because inherited variants in immune system genes increase susceptibility to certain infections. In support of this hypothesis, the risk that an adopted individual would die young from infection did not correlate with adoptive parents' death from infection before age 50. Although researchers concluded that length of life is mostly determined by heredity, they did find evidence of environmental influences. For example, if adoptive parents died before age 50 of cardiovascular disease, their adopted children were three times as likely to die of heart and blood vessel disease as a person in the general population. What environmental factor might explain this correlation?

Twins

Studies that use twins to separate the genetic from the environmental contribution to a phenotype provide more meaningful information than studying adopted individuals. Twin studies have largely replaced adoption methods. However, twin studies are not perfect experiments either. The genomes of identical twins are not really identical—they differ in DNA sequences called copy number variants (CNVs), which are repeats of short sequences. People differ in the numbers of repeats. CNVs are discussed further in chapter 12.

Using twins to study genetic influence on traits dates to 1924, when German dermatologist Hermann Siemens compared school transcripts of identical versus fraternal twins. Noticing that grades and teachers' comments were much more alike for identical twins than for fraternal twins, he proposed that genes contribute to intelligence.

A trait that occurs more frequently in both members of identical (monozygotic or MZ) twin pairs than in both members of fraternal (dizygotic or DZ) twin pairs is at least partly controlled by heredity. Geneticists calculate the **concordance** of a trait as the percentage of pairs in which both twins express the trait among pairs of twins in whom at least one has the trait. Twins who differ in a trait are said to be discordant for it. Copy number variant differences can explain some discordance among MZ twins.

In one study, 142 MZ twin pairs and 142 DZ twin pairs took a “distorted tunes test,” in which 26 familiar songs were played, each with at least one note altered. A person was considered “tune deaf” if he or she failed to detect the mistakes in three or more tunes. Concordance for “tune deafness” was

Table 7.5 Concordance Values for Some Traits in Twins

Trait	MZ (identical) twins	DZ (fraternal) twins
Acne	14%	14%
Alzheimer disease	78%	39%
Anorexia nervosa	55%	7%
Autism	90%	4.5%
Bipolar disorder	33–80%	0–8%
Cleft lip with or without cleft palate	40%	3–6%
Hypertension	62%	48%
Schizophrenia	40–50%	10%

67 percent for MZ twins, but only 44 percent for DZ twins, indicating a considerable inherited component in the ability to accurately perceive musical pitch. **Table 7.5** compares twin types for a variety of hard-to-measure traits. (Figure 3.16 shows how DZ and MZ twins arise.)

Diseases caused by single genes that approach 100 percent penetrance, whether dominant or recessive, also approach 100 percent concordance in MZ twins. That is, if one twin has the disease, so does the other. However, among DZ twins, concordance generally is 50 percent for a dominant trait and 25 percent for a recessive trait. These are the Mendelian values that apply to any two siblings. For a polygenic trait with little environmental input, concordance values for MZ twins are significantly greater than for DZ twins. A trait molded mostly by the environment exhibits similar concordance values for both types of twins.

Comparing twin types assumes that both types of twins share similar experiences. In fact, MZ twins are often closer emotionally than DZ twins. This discrepancy between the closeness of the two types of twins can lead to misleading results. A study from the 1940s, for example, concluded that tuberculosis is inherited because concordance among MZ twins was higher than among DZ twins. Actually, the infectious disease more readily passed between MZ twins because their parents kept them closer. However, the 1940s study wasn't totally off the mark. We do inherit susceptibilities to some infectious diseases. MZ twins would share such genes, whereas DZ twins would only be as likely as any sibling pairs to do so.

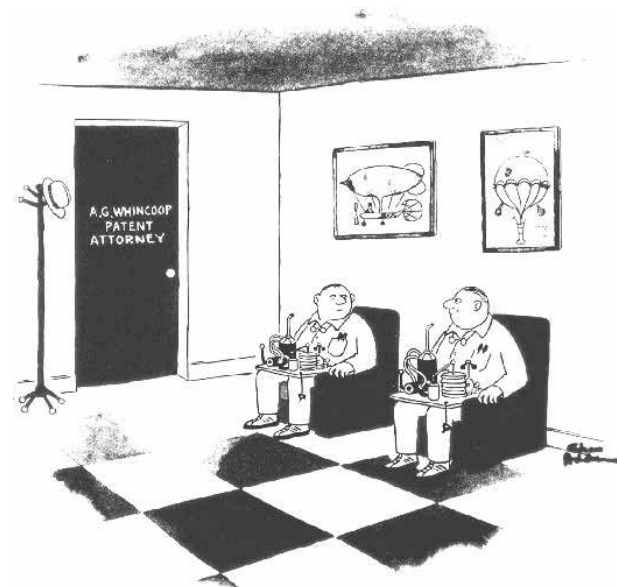
For some traits for which the abnormality may produce symptoms before birth, the type of MZ twin may be important. That is, MZ twins with the same amnion may share more environmental factors than MZ twins who have separate amnions (see figure 3.16). Schizophrenia is a condition that may begin subtly, before birth, and later become obvious when environmental factors come into play. Schizophrenia is discussed in chapter 8.

A more informative way to assess the genetic component of a multifactorial trait is to study MZ twins who were separated at birth, then raised in very different environments. Much of the work

using this “twins reared apart” approach has taken place at the University of Minnesota. Here, since 1979, hundreds of sets of twins and triplets who were separated at birth have visited the laboratories of Thomas Bouchard. For a week or more, the twins and triplets are tested for physical and behavioral traits, including 24 different blood types, handedness, direction of hair growth, fingerprint pattern, height, weight, functioning of all organ systems, intelligence, allergies, and dental patterns. Researchers videotape facial expressions and body movements in different circumstances and probe participants' fears, interests, and superstitions.

Twins and triplets separated at birth provide natural experiments for distinguishing nature from nurture. Many of their common traits can be attributed to genetics, especially if their environments have been very different. By contrast, their differences tend to reflect differences in upbringing, since their genes are identical (MZ twins and triplets) or similar (DZ twins and triplets).

MZ twins and triplets separated at birth and reunited later are remarkably similar, even when they grow up in very different adoptive families (**figure 7.8**). Idiosyncrasies are



Separated at birth, the Mallifert twins meet accidentally.

Figure 7.8 MZ twins separated at birth and reunited as adults may have astounding similarities. Originally published in the 4 May 1981 issue of *The New Yorker Magazine*, p. 43. © Tee and Charles Addams Foundation. Reprinted by permission.

particularly striking. One pair of twins who met for the first time when they were in their thirties responded identically to questions; each paused for 30 seconds, rotated a gold necklace she was wearing three times, and then answered the question. Coincidence, or genetics?

The “twins reared apart” approach is not an ideal way to separate nature from nurture. MZ twins and other multiples share an environment in the uterus and possibly in early infancy that may affect later development. Siblings, whether adoptive or biological, do not always share identical home environments. Differences in sex, general health, school and peer experiences, temperament, and personality affect each individual’s perception of such environmental influences as parental affection and discipline.

7.4 Genome-Wide Association Studies

The pedigrees in chapter 4 and linkage studies described in chapter 5 follow single-gene traits from generation to generation. The empiric risk, heritability, and adoptee and twin studies discussed in the first half of this chapter address conditions to which many genes contribute. Sequencing of the human genome and another large effort called the HapMap project, which identified SNPs, have led to a new tool, the genome-wide association study. In contrast to the older techniques that search for known gene variants, typically in only a few people, genome-wide association studies canvass signposts dispersed throughout the genome in many individuals to tease out the common variants behind a particular phenotype. An underlying assumption is that more common disorders with inherited causes remain with us because they affect health later in life, after a person has had children and passed on those gene variants.

The National Institutes of Health (NIH) defines a genome-wide association study as “any study of genetic variation across the entire human genome that is designed to identify genetic associations with observable traits (such as blood pressure or weight), or the presence or absence of a disease or condition” (figure 7.9). To achieve statistical significance, a genome-wide association study must include at least 100,000 markers, most of which are SNPs or copy number variants that are inherited. Alternatively, the studies may compare gene expression patterns. This approach can discover genes whose encoded proteins are over- or under-produced in a particular illness, such as proteins that are part of a general inflammatory response. Many media reports that “link” traits to “genes” are actually reporting associated gene expression patterns, not inheritance of mutations.

Recall from chapter 1 that a SNP (single nucleotide polymorphism) is a site in a genome where the DNA base varies in at least 1 percent of a population. “Polymorphism” simply means “variant.” Figure 7.10 shows three SNPs in part of a gene whose protein product enables a

person to taste very bitter substances. The three sites form two haplotypes, which function as alleles—a person is either a “taster” or a “non-taster.”

In genome-wide association studies, SNPs span the genome, rather than define a single gene. A SNP can be anywhere among our 3.2 billion base pairs—it does not have to be in a protein-encoding sequence. It is the association of SNP to trait that is informative. That is, if a SNP always occurs in individuals who share a specific trait, then it may do so because it lies in or near (linked to) a gene that *does* cause the trait—a genetic form of guilt-by-association.

Most investigations use a “tag SNP” that is inherited with others close to it on a chromosome. In this way, the millions of SNPs in the genome are grouped into 500,000 haplotypes. Following tag SNPs is a little like recognizing sports teams by identifying the captains.

Designing a Genome-Wide Association Study

A genome-wide association study is a step-wise focusing in on parts of the genome responsible to some degree for a trait (figure 7.11). In general, a population and a control group have their DNA isolated and genotyped for the 500,000 tag SNPs. Statistical algorithms identify the uniquely shared SNPs in the group with the trait or disorder of interest, and then the association is validated by repeating the process on additional populations. Each iteration narrows the SNPs and strengthens the association. It is important to validate a SNP association in different population groups, to be certain that it is the trait of interest that is being tracked, and not another part of the genome that members of one population share due to their common ancestry.

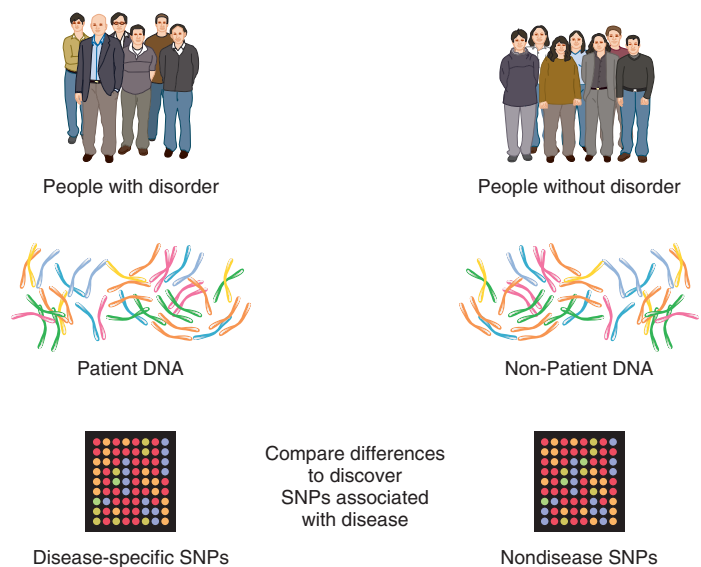


Figure 7.9 The new way to track genes. Genome-wide association studies seek DNA sequence variants that are shared with much greater frequency among individuals with the same illness or trait than among others.

PTC gene, non-taster allele

```

1      atgttgactc taactcgcac cgcactgtg tcctatgaag tcaggagtac atttctgttc
61..  atttcagtcc tggagtttgc agtggggttt ctgaccaatg ccttcgtttt cttggtgaat
121.. ttttgggatg tagtgaagag gcagcactg agcaacagtg attgtgtgct gctgtgtctc
...

781   tgtgttgcct tcactctctg gccctactg attctgtggc gcgacaaaat aggggtgatg
841   gtttgtgttg ggataatggc agcttgtccc tctgggcatg cagccatcct gatctcaggc
901   aatgccaagt tgaggagagc tgtgatgacc attctgtctc gggctcagag cagcctgaag
961   gtaagagccg accacaaggc agattcccgg acactgtgct ga

```

PTC gene, taster allele

```

1      atgttgactc taactcgcac cgcactgtg tcctatgaag tcaggagtac atttctgttc
61..  atttcagtcc tggagtttgc agtggggttt ctgaccaatg ccttcgtttt cttggtgaat
121.. ttttgggatg tagtgaagag gcagcactg agcaacagtg attgtgtgct gctgtgtctc
...

781   tgtgttgcct tcactctctg gccctactg attctgtggc gcgacaaaat aggggtgatg
841   gtttgtgttg ggataatggc agcttgtccc tctgggcatg cagccatcct gatctcaggc
901   aatgccaagt tgaggagagc tgtgatgacc attctgtctc gggctcagag cagcctgaag
961   gtaagagccg accacaaggc agattcccgg acactgtgct ga

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Figure 7.10 SNPs are sites of variability in genomes. These two sets of DNA base sequences represent a gene that controls the ability to taste phenylthiocarbamide (PTC). About 70 percent of individuals taste an extremely bitter substance when they lick a paper containing PTC; the other 30 percent taste nothing at all. Sequencing the PTC gene revealed differences at three sites. In the illustration, the numbers on the left are guidelines to position within the gene—the entire gene is 1,003 bases long. The bases are displayed in groups of ten for ease of counting. In a nontaster, positions 145, 785, and 886 are G, T, and A, respectively. In a taster, these three DNA bases are instead C, C, and G. The changes substitute three different amino acids in the encoded protein, which is sufficient to alter the phenotype.

Several different study designs are used in these investigations. In a **cohort study**, researchers follow a large group of individuals over time and measure many aspects of their health. The most famous is the Framingham Heart Study, which began tracking thousands of people and their descendants in

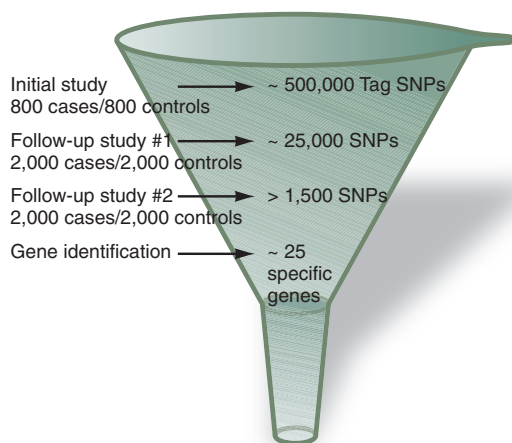


Figure 7.11 A stepwise approach to gene discovery. Genome-wide association study results must be validated in several different populations. Further research is necessary to go beyond association to demonstrate correlation and cause.

Massachusetts in 1968. Nine thousand of them are participating in a genome-wide association study. The gout investigation described in the chapter opening essay used the Framingham cohort along with two others, measuring uric acid levels as well as gout diagnoses to implicate three genes.

Genome-wide association studies can be set up in different ways. In a **case-control study**, each individual in one group is matched to an individual in another group who shares as many characteristics as possible, such as age, sex, activity level, and environmental exposures. SNP differences are then associated with the presence or absence of the disorder. For example, if 5,000 individuals with hypertension (high blood pressure) have particular DNA bases at six sites in the genome, and 5,000 matched individuals who do not have hypertension have different bases at only these six sites, then these genome regions may include genes whose protein products control blood pressure.

The “affected sibling pair” strategy borrows from the older technique of tracking linkage in families. Researchers scan genomes for SNPs that most siblings who have the same condition share, but that siblings who do not both have the condition do not often share. Such genome regions may harbor genes that contribute to the condition. The logic is that because siblings share 50 percent of their genes, a trait or condition that many siblings share is likely to be inherited.

A variation on the affected sibling pair strategy is **homozygosity mapping**, which is performed on families

that are consanguineous—that is, the parents are related. The genomes of children whose parents share ancestors have more homozygous regions than do other children, and therefore greater likelihood that they will inherit two copies of a susceptibility or disease-causing mutation. Homozygosity mapping was used to identify genes that cause autism, discussed in the next chapter.

Once a SNP association has been validated in diverse and large populations, gene identification can begin. This might entail consulting the human genome sequence for “candidate” genes near associated SNPs whose known functions make sense in explaining the condition, such as the three gout genes that affect the kidney’s excretion of uric acid. Or, understanding the SNP association might require exploring the phenotype. For example, a set of SNPs might track with breast cancer. Examination of the tumors at the molecular, cell, and tissue levels might reveal a subtype of the cancer that the SNP signature identifies. Then perhaps a test for the SNPs might be used to refine diagnosis.

Genome-wide association studies are also used to probe common characteristics, such as height. The results estimate the degree to which a particular gene variant contributes to the variation of a trait in a population. For example, a genome-wide association study of height identified variants in a gene called *HMGA2* as “explaining about 0.3 percent of population variation in height,” or about 0.4 centimeters of height per allele. In addition to all of the data, common sense comes into play. Inheriting several gene variants that each add to height is meaningless if a person doesn’t eat enough to grow.

Limitations of Genome-Wide Association Studies

Genome-wide association studies are prone to error simply because they include so many data points—but, ironically, it is the large numbers of markers, measurements, and people that build accuracy in pointing toward genes that affect health. To get an idea of the computational magnitude of such a study, consider a small version: An investigation of 20 genes, each with four SNPs. That equals 160 data points per individual ($20 \text{ genes} \times 4 \text{ SNPs/gene} \times 2 \text{ copies of each gene}$). With thousands of possible combinations (genotypes) of those 160 points, it’s clear that many thousands of individuals would have to be examined to note any SNP patterns that people who share symptoms have in common. Now imagine a study with a million SNPs, as is common today.

Another limitation of genome-wide association studies is that they reveal associations between two types of information, and not causes. An *association* only means that one event or characteristic occurs when another occurs. A *correlation* is a directional association: If one measurement increases, so does the other, such as stress and blood pressure. In contrast, establishing a cause requires that a specific mechanism explains how one event makes another happen: *How* does stress elevate blood pressure? An association study

does not provide information on a gene’s function—it is more a discovery tool.

How the patient population is selected can introduce bias into a genome-wide association study. Samples drawn from clinics, for example, would not include the very mildly affected or those who have died. It would also miss individuals for whom a contributing gene variant is nonpenetrant, or is associated with a late-onset disorder. Another source of error is that individuals in the control population might not actually be healthy—they might have problems other than the one being investigated.

The complicating factors discussed in chapter 5 also affect the accuracy of genome-wide association studies. Recall that a phenocopy is a trait or illness that resembles an inherited one, but has an environmental cause. Placing a person with anemia due to a drug reaction in a group with people who have an inherited anemia would be misleading. Genetic heterogeneity, in which different genes cause the same trait or condition, could also be a source of error. Epistasis, when one gene masks the effect of another, also confounds these studies, but as we learn more, these interactions are being taken into account.

Yet another source of error arises from what a genome-wide association study misses, such as extremely rare SNPs. Or, the people who share symptoms and a SNP pattern may share something else that accounts for the association, such as an environmental exposure. This situation can generate a false positive result.

Many of these limitations are overcome by expanding the numbers, and pooling data. Indeed, when these studies began in the early part of the century, associations would often fall apart as the numbers grew, and it was not uncommon to see results reported one year retracted the next. Still, some researchers think that these limitations compromise the value of genome-wide association studies. This is especially true for cases in which heritability calculated using more traditional approaches is much higher than the genetic contribution revealed in the study. However, the fact that genome-wide association studies do not uncover all of the genetic contribution to a trait or illness may just mean that we haven’t yet found all the contributing genes.

Often, the old and the new techniques for dissecting multifactorial traits work well together. This is the case for stuttering. Concordance for MZ twins ranges from 20 to 83 percent, and for DZ twins, from 4 to 9 percent, suggesting a large inherited component. The risk of a first-degree relative of a person who stutters also stuttering is 15 percent based on empiric evidence, compared to the lifetime risk of stuttering in the general population of 5 percent, although part of that increase could be due to imitating an affected relative. A genome-wide association study on 100 families who have at least two members who stutter identified candidate genes on three chromosomes that contribute to the trait.

Genome-wide association studies have already examined dozens of conditions, and identified hundreds of genes, giving drug developers many new targets to work with, and explaining

Table 7.6 Genome-Wide Association Studies
in Progress(For details see <http://www.genome.gov/26525384>)

Autoimmune/Allergic/ Infectious disorders	AIDS progression	
	Asthma	
	Crohn's disease	
	Diabetes mellitus	
	Multiple sclerosis	
	Psoriasis	
	Rheumatoid arthritis	
	Systemic lupus erythematosus	
	Behavioral traits and disorders	Attention deficit hyperactivity disorder
		Autism
		Bipolar disorder
		Conduct disorder
		General cognitive ability
Major depressive disorder		
Panic disorder		
Personality		
Schizophrenia		
Sleep characteristics		
Smoking behavior		
Cancers	Breast	
	Blood	
	Colon	
	Prostate	
	Ovarian	
	Thyroid	
Cardiovascular disease	Coronary artery disease	
	Hypertension	
	Long QT syndrome	
	Myocardial infarction	
Neurological disorders	Alzheimer disease	
	Amyotrophic lateral sclerosis	
	Narcolepsy	
	Parkinson disease	
	Restless legs syndrome	
Other	Bone mineral density	
	Chronic obstructive pulmonary disease	
	Cleft lip/cleft palate	
	Drug responses	
	Gout	
	Male pattern baldness	
Physical traits	Body mass index	
	Brain volume	
	Eye color	
	Freckles	
	Hair color	
	Height	
	Facial feature spacing	
	Head circumference	
	Limb lengths	

how certain conditions arise. **Table 7.6** lists some of the conditions. **Table 7.7** reviews terms used to study multifactorial traits. The next section probes an example of such a trait—body weight.

Table 7.7 Terms Used in Evaluating
Multifactorial Traits

Coefficient of relatedness The proportion of genes shared by two people related in a particular way. Used to calculate heritability.

Concordance The percentage of twin pairs in which both twins express a trait.

Empiric risk The risk of recurrence of a trait or illness based on known incidence in a particular population.

Genome-wide association study Detecting association between marker patterns and increased risk of developing a particular medical condition.

Heritability The percentage of phenotypic variation for a trait that is attributable to genetic differences. It equals the ratio of the observed phenotypic variation to the expected phenotypic variation for a population of individuals.

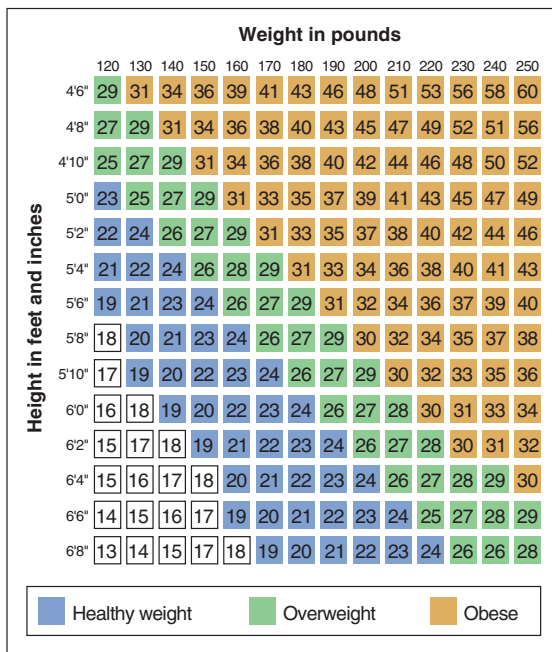
Key Concepts

1. Researchers compare traits in adopted individuals to those in their adoptive and biological parents to assess the genetic contribution to a trait.
2. Concordance is the percentage of twin pairs in which both express a trait. For a trait largely determined by genes, concordance is higher for MZ than DZ twins.
3. Genome-wide association studies seek correlations between SNP patterns and phenotypes in large groups of individuals.

7.5 A Closer Look: Body Weight

Unlike rare genetic disorders, body weight is a multifactorial trait that we all have. Body weight reflects energy balance—the rate of food taken in versus the rate at which the body uses it for fuel. Excess food means, ultimately, excess weight. About 30 percent of all adults in the United States are obese, and another 35 percent are overweight. Being overweight or obese raises the risk of developing hypertension, diabetes, stroke, gallstones, sleep apnea, and some cancers.

Scientific studies of body weight use a measurement called body mass index (BMI), which is weight in proportion to height (**figure 7.12**). BMI makes sense—a person who weighs 170 pounds and is 6 feet tall is slim, whereas a person of the same weight who is 5 feet tall is obese. The tall person's BMI is 23; the short person's is 33.5.



Developed by the National Center for Health Statistics in collaboration with the National Center for Chronic Disease Prevention and Health Promotion

Figure 7.12 Body mass index (BMI). BMI equals weight/height², with weight measured in kilograms and height measured in meters. This chart provides a shortcut—the calculations have been done and converted to the English system of measurement. Squares that are not filled in indicate underweight.

Heritability for BMI is 0.55, which leaves room for environmental influences on our appetites and sizes. Some genes implicated in determining body weight have been known for a long time. New ones, some with unexpected functions, have shown up in genome-wide association studies, which point to more than 50 genome regions that may harbor genes that affect how much we eat, how we use calories, and how fat is

distributed in the body. The biochemical pathways and hormonal interactions that control weight may reveal points for drug intervention (table 7.8).

Leptin and Associated Proteins

Obesity research first embraced genetics in 1994, when Jeffrey Friedman at Rockefeller University discovered a gene that encodes the protein hormone leptin in mice and in humans. Normally, eating stimulates fat cells (adipocytes) to secrete leptin, which travels in the bloodstream to a region of the brain’s hypothalamus, where it binds to receptors on nerve cells (neurons). The binding signals the neurons to release another type of hormone that binds yet other types of receptors, which ultimately function as an appetite “brake,” while speeding breakdown of food already eaten. When a person hasn’t eaten in several hours, leptin levels ebb, which triggers the release of an appetite “accelerator.” Table 7.8 lists the details of some proteins that affect eating behavior.

The discovery of genes and proteins that affect appetite led to great interest in targeting them with drugs to either lose or gain weight. When Friedman gave mice extra leptin, they ate less and lost weight. Headlines soon proclaimed the new magic weight loss elixir, a biotech company paid \$20 million for rights to the hormone, and clinical trials ensued. The idea was to give obese people leptin, assuming that they had a deficiency, to trick them into feeling full. Only about 15 percent of the people lost weight, but the other 85 percent didn’t actually lack leptin. Instead, most of them had leptin resistance, which is a diminished ability to recognize the hormone due to defective leptin receptors. Giving these people leptin had no effect on their appetites. However, the discovery helped a few severely obese children with true leptin deficiency attain normal weights after years of daily leptin injections.

The stomach is another source of obesity-related proteins. Ghrelin is a peptide (small protein) hormone produced in the stomach that responds to hunger, signaling the hypothalamus

Table 7.8 Some Sites of Genetic Control of Body Weight

Protein	Function	MIM	Effect on Appetite
Leptin	Stimulates cells in hypothalamus to decrease appetite and metabolize nutrients.	164160	↓
Leptin transporter	Enables leptin to cross from bloodstream into brain.	601694	↓
Leptin receptor	Binds leptin on hypothalamus cell surfaces, triggering hormone’s effects.	601007	↓
Neuropeptide Y	Produced in hypothalamus when leptin levels are low and the individual loses weight.	162640	↑
Melanocortin-4 receptor	Activated when leptin levels are high and the individual gains weight.	155541	↓
Ghrelin	Signals hunger from stomach to brain in short term, stimulating neuropeptide Y.	605353	↑
PYY	Signals satiety from stomach to brain.	660781	↓
Stearoyl-CoA desaturase-1	Controls whether body stores or uses fat.	604031	↑

to produce more of the appetite accelerator. A different peptide hormone opposes ghrelin, signaling satiety to the brain. A drug is being developed to block the effects of ghrelin, which may one day replace weight-loss surgery with a pill. While leptin acts in the long term to maintain weight, the stomach's appetite control hormones function in the short term. All of these hormonal signals are integrated to finely control appetite in a way that maintains weight.

Beyond Single Weight-Control Genes

Identifying single genes that influence weight paved the way for considering the trait to be multifactorial. Researchers are investigating combinations of known genes as well as many newly discovered genes.

One study looked at 21 genes in which mutations cause syndromes that include obesity, as well as 37 genes whose products participate in biochemical pathways related to weight. This approach identified many rare gene variants that could, in combinations, explain many people's tendency to gain weight. In another study of known genes, the effectiveness of a weight loss drug (Meridia) was associated with certain variants of the genes that encode the receptors that two appetite-controlling neurotransmitters bind.

Genome-wide association studies that compare gene expression patterns have also enhanced understanding of body weight. One study compared the sets of genes that are expressed in adipose (fat) tissue to other tissues. Samples from more than 1,600 people in Iceland revealed a set of genes whose products take part in inflammation and the immune response, but also contribute obesity-related traits. This was not known, and can suggest new drug targets.

Environmental Influences on Weight

Many studies on adopted individuals and twins suggest that obesity has a heritability of 75 percent. Because the heritability for BMI is lower than this, the discrepancy suggests that genes play a larger role in those who tend to gain weight easily. This becomes obvious when populations that have a genetic tendency to obesity experience a large and sudden change in diet for the worse.

On the tiny island of Nauru, in Western Samoa, the residents' lifestyles changed greatly when they found a market for the tons of bird droppings on their island as commercial fertilizer. The influx of money translated into inactivity and a high-calorie, high-fat diet, replacing an agricultural lifestyle and a diet of fish and vegetables. Within just a generation, two-thirds of the population had become obese, and a third suffered from diabetes.

The Pima Indians offer another example of environmental effects on body weight. These people separated into two populations during the Middle Ages, one group settling in the Sierra Madre mountains of Mexico, the other in southern Arizona. By the 1970s, the Arizona Indians no longer farmed nor

ate a low-calorie, low-fat diet, but instead consumed 40 percent of their calories from fat. With this extreme change in lifestyle, they developed the highest prevalence of obesity of any population on earth. Half of the Arizona group had diabetes by age 35, weighing, on average, 57 pounds (26 kilograms) more than their southern relatives, who still eat a low-fat diet and are very active.

The Pima Indians demonstrate that future obesity is not sealed in the genes at conception, but instead is much more likely to occur if the environment provides too many calories and too much fat. They illustrate what geneticist James Neel termed the "thrifty gene hypothesis" in 1962. He suggested that long ago, the hunter-gatherers who survived famine had genes that enabled them to efficiently conserve fat. Today, with food plentiful, the genetic tendency to retain fat is no longer healthful, but harmful. Unfortunately, for many of us, our genomes hold an energy-conserving legacy that works too well—it is much easier to gain weight than to lose it, for sound evolutionary reasons.

The thrifty gene hypothesis is also seen in people who were born after a full-term pregnancy, but were very low weight. To compensate for starvation conditions in the uterus, metabolism shifts in a way that conserves calories—and the person pays for it later with elevated risk of heart disease, stroke, obesity, osteoporosis, and type 2 diabetes. These are multifactorial conditions that, instead of arising from mutations, reflect epigenetic alterations of gene expression.

Another environmental influence on weight is the types of bacteria in our bodies. Bacterial cells in our bodies actually outnumber our own cells. The actions of certain types of bacteria affect the number of calories that we extract from particular foods. Researchers demonstrated this by controlling the diets of a group of obese individuals and monitoring the bacterial species in their feces. The investigators identified nine species of bacteria that enable a human body to extract maximal calories from food.

Perhaps nowhere are the complexities and challenges of gene-environment interactions more profound than in behavioral characteristics, nuances, quirks, and illnesses. The next chapter looks at a few of them.

Key Concepts

1. Genes that encode leptin, the leptin receptor, and proteins that transmit or counter leptin's signals affect body weight.
2. Studies on adopted individuals and twins indicate a heritability of 75 percent for obesity.
3. Populations that suddenly become sedentary and switch to a fatty diet reflect environmental influences on body weight.

Summary

7.1 Genes and the Environment Mold Traits

1. **Multifactorial traits** reflect the environment and genes. A **polygenic trait** is determined by more than one gene and varies continuously in expression.
2. Single-gene traits are rare. For most traits, many genes contribute to a small, but not equal, degree.

7.2 Polygenic Traits Are Continuously Varying

3. The frequency distribution of phenotypes for a polygenic trait forms a bell curve.

7.3 Traditional Approaches to Investigating Multifactorial Traits

4. **Empiric risk** measures the likelihood that a multifactorial trait will recur based on prevalence. The risk rises with genetic closeness, severity, and number of affected relatives.
5. **Heritability** estimates the proportion of variation in a multifactorial trait due to genetics in a particular population at a particular time.
6. Characteristics shared by adopted people and their biological parents are mostly inherited, whereas similarities

between adopted people and their adoptive parents reflect environmental influences.

7. **Concordance** measures the frequency of expression of a trait in both members of MZ or DZ twin pairs. The more influence genes exert over a trait, the higher the differences in concordance between MZ and DZ twins.

7.4 Genome-Wide Association Studies

8. **Genome-wide association studies** correlate SNP patterns to increased disease risk. They may use a **cohort study** to follow a large group over time, or a **case-control** study on matched pairs. The affected sibling pair strategy can identify homozygous regions that may harbor genes of interest.

7.5 A Closer Look: Body Weight

9. Many genes affect weight. Leptin and associated proteins affect appetite. Fat cells secrete leptin in response to eating, which acts in the hypothalamus to decrease appetite. Populations that switch to a fatty, high-calorie diet and a less-active lifestyle reveal effects of the environment on weight.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.

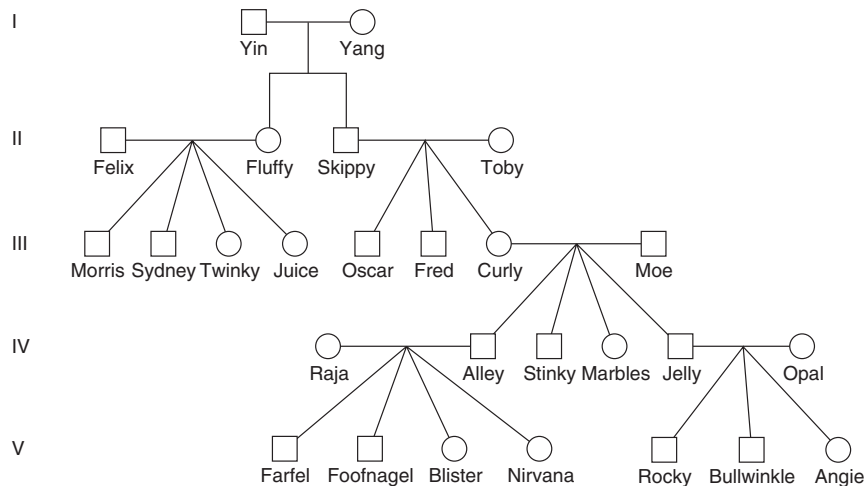


Review Questions

1. Explain how Mendel's laws apply to multifactorial traits.
2. Explain the difference in the genetic contribution to sickle cell disease or cystic fibrosis, compared to that for hypertension not due to another illness.
3. Why is recurrence of a disease that would fall on the lower right of figure 7.1 more difficult to predict than a disease depicted on the upper left?
4. What is the difference between a Mendelian multifactorial trait and a polygenic multifactorial trait?
5. How can skin color have a different heritability at different times of the year?
6. Explain how the twins in figure 7.4 have such different skin colors.
7. In a large, diverse population, why are medium brown skin colors more common than very white or very black skin?
8. Explain how the Connecticut and Northern Finland investigations of height differ in how they were conducted and in the type of information revealed.
9. Describe the type of information in a(n)
 - a. empiric risk calculation.
 - b. twin study.
 - c. adoption study.
 - d. genome-wide association study.
10. Which has a greater heritability—eye color or height? State a reason for your answer.
11. Why does SNP mapping require extensive data?
12. How can older techniques to study multifactorial traits be combined with newer techniques?
13. How can a genome-wide association study overcome the bias of looking only at genes already known to be involved in a trait, or whose known function makes them likely candidates?
14. Name three types of proteins that affect cardiovascular functioning and three that affect body weight.

Applied Questions

- Rebecca breeds Maine coon cats. The partial pedigree below describes how her current cats are related—the umbrellalike lines indicate littermates, which are the equivalent of fraternal (DZ) twins in humans.
Cat lover Sam wishes to purchase a pair of Rebecca's cats to breed, but wants them to share as few genes as possible to minimize the risk that their kittens will inherit certain multifactorial disorders. Sam is quite taken with Farfel, but can't decide among Marbles, Juice, or Angie for Farfel's mate.
Calculate the percentage of genes that Farfel shares with each of these female relatives. With which partner would the likelihood of healthy kittens be greatest?
- Cite an example from chapter 5 of a single-gene trait or condition that is affected by an environmental influence.
- Would you take a drug that was prescribed to you based on your race? Cite a reason for your answer.
- The incidence of obesity in the United States has doubled over the past two decades. Is this due more to genetic or environmental factors? Cite a reason for your answer.
- One way to calculate heritability is to double the difference between the concordance values for MZ versus DZ twins. For multiple sclerosis, concordance for MZ twins is 30 percent, and for DZ twins, 3 percent. What is the heritability? What does the heritability suggest about the relative contributions of genes and the environment in causing MS?
- Devise a genome-wide association study to assess whether restless legs syndrome is inherited, and if it is, where susceptibility or causative genes may be located.
- In chickens, high body weight is a multifactorial trait. Heritability accounts for several genes that contribute a small effect additively, as well as a few genes that exert a great effect. Is this an example of narrow or broad heritability?



Maine coon cats

Web Activities

- Genome-wide association studies are research tools used to reveal the genetic underpinnings of traits and diseases in populations. Some direct-to-consumer genetic testing websites offer "educational services" that report genetic marker patterns found in these studies that are associated with a particular disease—applied to the individual who purchases the test. Do you think this is a valid service? State a reason for your answer.
- Locate a website that deals with breeding show animals, farm animals, or crops to produce specific traits, such as litter size, degree of meat marbling, milk yield, or fruit ripening rate. Identify three traits with heritabilities that indicate a greater contribution from genes than from the environment.
- Visit the Centers for Disease Control and Prevention (CDC) website. From the leading causes of death, list three that have high heritabilities, and three that do not. Base your decisions on common sense or data, and explain your selections.
- Use MIM to look up any of the following genes that affect cardiovascular health and explain what the genes do: apolipoprotein E; LDL receptor; apolipoprotein A; angiotensinogen; beta-2 adrenergic receptor; toll-like receptor 4; C reactive protein.

Case Studies and Research Results

12. Concordance for the eating disorder anorexia nervosa for MZ twins is 55 percent, and for DZ twins, 7 percent. Ashley and Maggie are DZ twins. Maggie has anorexia nervosa. Should Ashley worry about an inherited tendency to develop the condition? Explain your answer.
13. Lydia and Reggie grew up poor in New York City in the 1960s. Both went for free to the City University of New York, then to medical school in Boston, where they met. Today, each has a thriving medical practice, and they are the parents of 18-year-old Jamal and 20-year-old Tanya.

Jamal, taking a genetics class, wonders why he and Tanya do not resemble each other, or their parents, for some traits. The family is African American. Lydia and Reggie are short, 5'2" and 5'7" respectively, and each has medium brown eyes and skin, and dark brown hair. Tanya and Jamal are 5'8" and 6'1", respectively, and were often in the highest height percentiles since they were toddlers. Jamal has very dark skin, darker than his parents' skin, while Tanya's skin is noticeably lighter than that of either parent. Tanya's eyes are so dark that they appear nearly black.

 - a. Why do Tanya's eyes appear darker than those of her parents or brother?
 - b. How can Jamal's skin be darker than that of his parents, and Tanya's be lighter?
 - c. Which of the traits considered is most influenced by environmental factors?
 - d. What is the evidence that Jamal and Tanya's height is due to environmental and genetic factors?
 - e. Which of the four traits has the highest heritability?
14. A study looked at 200,000 SNPs throughout the genome for 1,820 people with premature hair graying and 1,820 without this trait. Those with the trait shared several SNPs on chromosome 9. What type of study is this?
15. "Elite controllers" are people with HIV infection whose immune systems naturally keep levels of the virus extremely low. Researchers are conducting genome-wide SNP analyses of these people. Explain how the results of this study might be used to help people who more easily develop AIDS.
16. An affected sibling pair study identified areas of chromosomes 1, 14, and 20 that are likely to harbor genes that predispose individuals toward or cause schizophrenia. Explain how such an investigation is conducted.
17. Researchers compared the expression levels of 9,156 genes that could affect drug metabolism among 60 mother-father-child trios from Utah who were Caucasian, and 30 similar families who were black and from Nigeria. About 5 percent of the genes differed significantly in expression between the two groups. What further information would be helpful in applying this information in prescribing drugs?
18. A study is examining the expression of certain genes in people about to undergo weight loss surgery (gastric bypass), to see if these patterns predict individuals most likely to benefit by achieving long-term weight loss.
 - a. Name three genes, or types of genes, that might provide valuable information for this analysis.
 - b. What is a limitation of this study design?
 - c. Do you think that this study has value?



Chronic fatigue syndrome is not "all in one's head." It is likely due to an inherited extreme response to stress.

CHAPTER

8

Genetics of Behavior

Chronic Fatigue Syndrome

Chronic fatigue syndrome (CFS) usually begins with a flu-like illness, lasts at least six months, and causes disabling fatigue. Until recently, many people did not believe CFS was real. Genetics research suggests that it is.

Laura Hillenbrand, author of the bestseller *Seabiscuit*, has written about CFS. Ill since 1987, Laura has been bedbound for months on end, too exhausted to do anything. The disease forced her to drop out of college, and she lost a great deal of weight. Doctors blamed infection, and sent her to a psychiatrist, who ruled out mental illness. Until she finally found a physician who knew that CFS was real, Hillenbrand's doctors told her the disabling symptoms were "all in your head." They weren't.

Clues to the cause of CFS lay in the observation that many people report a severe physical illness or emotional trauma before the fatigue began. Also, people with CFS make too little of the stress hormone cortisol, and too much of the nervous system chemical serotonin, which induces sleep and calms mood. Could an abnormal and persistent response to a trauma—physical or emotional—cause CFS? Hormones from the brain's hypothalamus and pituitary gland, and the adrenal glands, control responses to trauma. Searching for variants of genes whose encoded proteins affect these hormones led to the discovery of three genes with variants that differ in people with CFS. One gene encodes the receptor that binds stress hormones and the other two affect the availability of serotonin. A physical explanation for CFS may lead to treatment.

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- 8.1 Genes and Behavior**
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 - Major Depressive Disorder
 - Bipolar Disorder
- 8.7 Schizophrenia**
 - Signs and Symptoms
 - Genetic Associations
 - Environmental Influences
- 8.8 Autism**

8.1 Genes and Behavior

Behavior is a complex continuum of emotions, moods, intelligence, and personality that drives how we function on a daily basis. We are, to an extent, defined and judged by our many behaviors. They control how we communicate, cope with negative feelings, and react to stress. At the end of the continuum of our behaviors are several common disorders, with wide-ranging and sometimes overlapping symptoms. Many aspects of our behavior occur in response to environmental factors, but *how* we respond has genetic underpinnings. Understanding the biology behind behavior can help to develop treatments for behavior-based disorders. Several are discussed in this chapter.

Behavioral genetics considers nervous system function and variation, including the hard-to-define qualities of mood and mind. The human brain weighs about 3 pounds and resembles a giant gray walnut, but with the appearance and consistency of pudding. It consists of 100 billion nerve cells, or neurons, and at least a trillion other cells called neuroglia, which support and nurture the neurons.

Connectivity and interaction among neurons make the brain a coordinated, functioning organ that controls all of the body. Branches from each of the 100 billion neurons in the brain

form close associations, called synapses, with 1,000 to 10,000 other neurons. Neurons communicate across these tiny spaces using chemical signals called neurotransmitters. The neurons form networks and clusters that oversee broad functions such as sensation and perception, memory, reasoning, and muscular movements.

Genes control the production and distribution of neurotransmitters. **Figure 8.1** indicates the points where genes control the sending and receiving of nervous system information. Enzymes oversee the synthesis of neurotransmitters and their transport from the sending (presynaptic) neuron across the synapse to receptors on the receiving (postsynaptic) neuron. Proteins called transporters ferry neurotransmitters from sending to receiving neurons, and proteins also form the subunits of receptors. Genes also control the synthesis of myelin, a fatty substance that coats neuron extensions called axons. Myelin coats and insulates the neuron, which speeds neurotransmission. Signal transduction is also a key part of the function of the nervous system (see figure 2.20). Therefore, candidate genes for the inherited components of a variety of mood disorders and mental illnesses—as well as of normal variations in temperament and personality—affect neurotransmission and signal transduction.

In contrast to investigating candidate genes selected by what is known about nerve transmission and signal transduction are genome-wide association studies, discussed in chapter 7. These investigations can identify genes not already suspected of affecting behavior.

Identifying the inherited and environmental contributors to a behavioral disorder is very challenging, partly because of the traditional way that psychiatrists diagnose such conditions. The widely used *Diagnostic and Statistical Manual (DSM)* categorizes and distinguishes mental and behavioral conditions based on symptoms. However, different syndromes share symptoms, and the same disorder can have different causes in different people.

Genetic studies are challenging the long-standing DSM classifications. Behavioral traits and disorders, like other characteristics, might reflect a major influence from a single gene, perhaps one whose protein product takes part directly in neurotransmission, but also small inputs from common gene variants. Researchers envision from 100 to 300 genes at play. Inheriting certain subsets of variants of these genes makes an individual susceptible to developing a certain

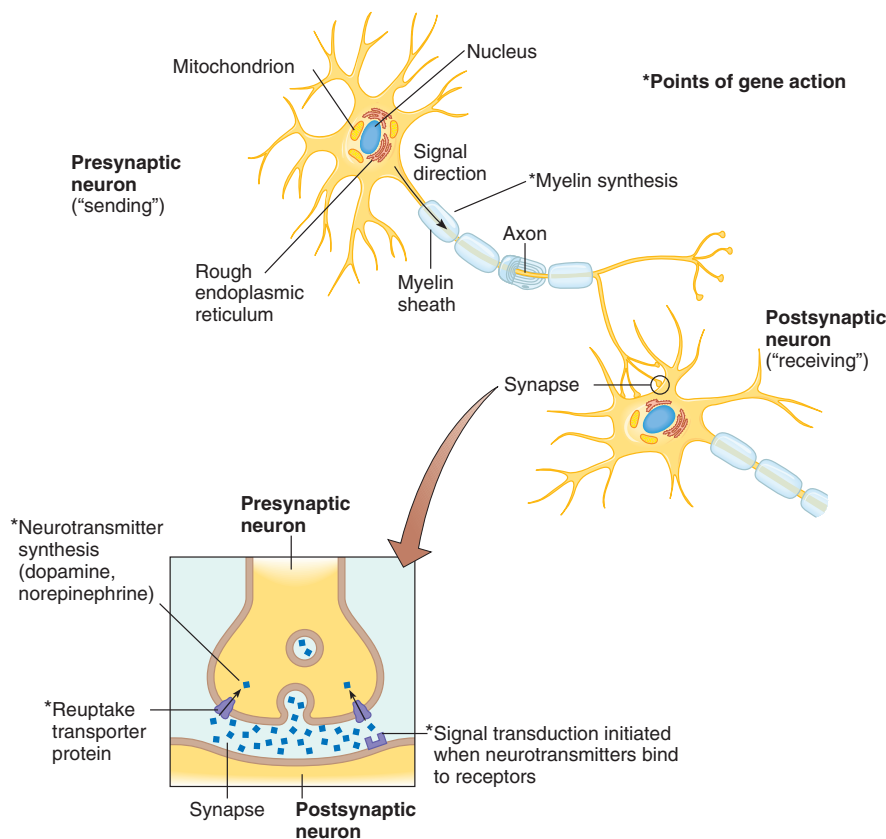


Figure 8.1 Neurotransmission. Many genes that affect behavior produce proteins that affect neurotransmission, which is sending a signal from one neuron to another across a synapse via a neurotransmitter molecule.

disorder in the presence of a particular environmental stimulus. Some genes are implicated in more than one behavioral disorder.

Rather than the separate designations in the phenotype-based DSM, genetics is showing that behavioral and mental disorders lie more across a continual spectrum. The Psychiatric Genome-Wide Association Study Consortium, for example, is one effort that is probing the complex genetic combinations that increase risk for behavioral disorders. In this project, investigators from Europe and North America are looking among 59,000 people for SNP patterns associated with autism, attention deficit hyperactivity disorder, bipolar disorder, depression, and schizophrenia. Another study is examining copy number variants (repeated short DNA sequences) that may explain behavioral disorders that occur sporadically, arising in someone without a family history.

Deciphering the genetic components of behavioral traits and disorders uses traditional empiric risk, adoptee, and twin study data and, more and more, molecular methods that determine genotypes, gene expression patterns, and sets of susceptibility gene variants. The chapter sections that follow explore these various sources of data, some beginning with case histories. Identifying gene variant combinations behind behavioral disorders may make it possible to subtype these conditions in a biologically meaningful way so that diagnoses will be more personalized and effective treatments begun sooner.

Table 8.1 lists the prevalence of some behavioral disorders.

Condition	Prevalence (%)
Alzheimer disease	4.0
Anxiety	8.0
Phobias	2.5
Posttraumatic stress disorder	1.8
Generalized anxiety disorder	1.5
Obsessive compulsive disorder	1.2
Panic disorder	1.0
Attention deficit hyperactivity disorder	2.0
Autism spectrum disorders	0.1
Drug addiction	4.0
Eating disorders	3.0
Mood disorders	7.0
Major depressive disorder	6.0
Bipolar disorder	1.0
Schizophrenia	1.3

Key Concepts

1. Behavioral traits reflect genetic and environmental influences, and arise from connections among the brain's 100 billion neurons.
2. The genes behind behavior affect neurotransmitter production and transport across synapses; myelin synthesis; and signal transduction.
3. Genetic classification of behavioral disorders is challenging traditional psychiatric classification. Behavioral disorders may lie on a continuum, with many genes having input.

8.2 Eating Disorders

In anorexia nervosa, a person perceives herself or himself as obese, even when obviously not, and intentionally starves, exercises obsessively, and/or takes laxatives to hasten weight loss (**figure 8.2**).

For economically advantaged females in the United States, the lifetime risk of developing anorexia nervosa is 0.5 percent. Anorexia has the highest risk of death of any psychiatric disorder—15 to 21 percent. The same population group has a lifetime risk of 2.5 percent of developing another eating disorder, bulimia. A person with bulimia eats huge amounts but exercises and vomits to maintain weight.

Five to 10 million people in the United States have eating disorders. About 10 percent of them are male. One survey of 8-year-old boys revealed that more than a third of them had attempted to lose weight. In an eating disorder called muscle dysmorphia, boys and young men take amino acid supplements



Figure 8.2 An eating disorder. In anorexia nervosa, a person perceives herself as obese although she is the opposite.

to bulk up. Just as the person with anorexia looks in a mirror and sees herself as too large, a person with muscle dysmorphia sees himself as too small.

Because eating disorders were once associated almost exclusively with females, most available risk estimates exclude males. Twin studies reveal a considerable genetic component, with heritability ranging from 0.5 to 0.8. Studies of eating disorders that recur in siblings can be difficult to interpret. Is a young girl imitating her older sister by starving because genes predispose her to develop an eating disorder, or because she wants to be like her sister?

Genes that encode proteins that control appetite are candidate genes for developing eating disorders (see table 7.8). Japanese researchers have performed a case-control, genome-wide association study for anorexia nervosa and identified four chromosomal regions that include genes that are active in the part of the brain that regulates eating behavior. It will be interesting to learn which genes affect body image, and how they do so.

Key Concepts

1. Eating disorders are common and have high heritability.
2. Genes whose products control appetite or regulate certain neurotransmitters may elevate the risk of developing an eating disorder.

8.3 Sleep

Sleep has been called “a vital behavior of unknown function,” and, indeed, without sleep, animals die. We spend a third of our lives in this mysterious state.

Genes influence sleep characteristics. When asked about sleep duration, schedule, quality, nap habits, and whether they are “night owls” or “morning people,” MZ twins report significantly more in common than do DZ twins. This is true even for MZ twins separated at birth. Twin studies of brain wave patterns through four of the five stages of sleep confirm a hereditary influence. The fifth stage, REM sleep, is associated with dreaming and therefore may reflect the input of experience more than genes.

Narcolepsy

Researchers discovered the first gene related to sleep in 1999, for a condition called “narcolepsy with cataplexy” in dogs. Humans have the disorder (MIM 161400), but it is rarely inherited as a single-gene trait—it is more often polygenic requiring an environmental trigger.

A person (or dog) with narcolepsy falls asleep suddenly several times a day. Extreme daytime sleepiness greatly disrupts daily activities. People with narcolepsy have a tenfold higher rate of car accidents. Another symptom is sleep

paralysis, which is the inability to move for a few minutes after awakening. The most dramatic manifestation of narcolepsy is cataplexy. During these short and sudden episodes of muscle weakness, the jaw sags, the head drops, knees buckle, and the person falls to the ground. This often occurs during a bout of laughter or excitement—which can be quite disturbing both for the affected individual and bystanders. People with narcolepsy and cataplexy cannot participate in even the most mundane of activities for fear of falling and injuring themselves. Narcolepsy with cataplexy affects only 0.02 to 0.06 percent of the general populations of North America and Europe, but the fact that it is much more common in certain families suggests a genetic component.

Studies on dogs led the way to discovery of a human narcolepsy gene. In 1999, Emmanuel Mignot and his team at Stanford University identified mutations in a gene that encodes a receptor for a neuropeptide called hypocretin (MIM 602358). In Doberman pinschers and Labrador retrievers, the receptor does not reach the cell surfaces of certain brain cells, and as a result, the cells cannot receive signals to promote a state of awakesness. Dachshunds have their own mutation—they make a misshapen, nonfunctional receptor. **Figure 8.3** shows a still frame of a film that Mignot made of narcoleptic dogs playing. Suddenly, they all collapse! A minute later, they get up and resume their antics. “You can’t make dogs laugh, but you can make them so happy that they have attacks,” says Mignot. To induce a narcoleptic episode in puppies, he lets them play with each other. He feeds older dogs meat, which excites them so much that they can take a while to finish a meal because they fall down in delight so often. Getting narcoleptic dogs to breed is difficult, too, for sex is even more exciting than play or food!

In 1998, Masahi Yanagisawa, at the University of Texas Southwestern Medical Center in Dallas, discovered a protein called orexin, but thought it only sent signals to eat. Yanagisawa’s orexin turned out to bind Mignot’s hypocretin receptor.



Figure 8.3 Letting sleeping dogs lie. These Doberman pinschers have inherited narcolepsy. They suddenly fall into a short but deep sleep while playing. Research on dogs with narcolepsy led to the discovery of the gene in humans.

Yanagisawa bred mice that lacked the orexin gene, and then noticed something odd while watching the animals feed at night—the rodents suddenly fell down fast asleep! Researchers are now trying to figure out how one molecule controls feeding as well as wakefulness. The hypocretin/orexin receptor gene, found on dog chromosome 12, is on human chromosome 6. The brains of humans with narcolepsy and cataplexy are remarkably deficient in hypocretin/orexin. A narcolepsy drug might mimic the missing molecule. Genome-wide association studies have identified other genes whose expression is greatly lowered in people with narcolepsy compared to those in controls.

Familial Advanced Sleep Phase Syndrome

Daily rhythms such as the sleep-wake cycle are set by cells that form a “circadian pacemaker” in two clusters of neurons in the brain called the suprachiasmatic nuclei. In these cells, certain “clock” genes are expressed in response to light or dark in the environment.

The function of clock genes is most obvious in families that have a mutation. For example, five generations of a family in Utah have familial advanced sleep phase syndrome (MIM 604348). Affected individuals promptly fall asleep at 7:30 each night and awaken suddenly at 4:30 A.M., thanks to a mutation in a gene on chromosome 2 called *period* (figure 8.4). The gene has counterparts in golden hamsters and fruit flies that disrupt their sleep-wake cycles, too. People with the condition have a single DNA base substitution that prevents the encoded protein from binding a phosphate chemical group, which it must do to signal to the brain in a way that synchronizes the sleep-wake cycle with daily sunrise and sunset. At least three other genes are associated with insomnia.

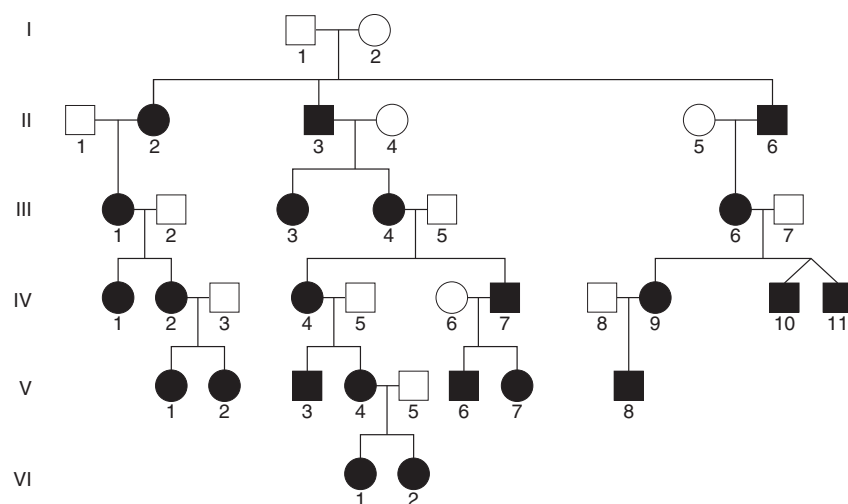


Figure 8.4 Inheritance of a disrupted sleep-wake cycle. This partial pedigree depicts affected members of a large family with an autosomal dominant form of familial advanced sleep phase syndrome.

Key Concepts

1. Twin studies on sleep habits indicate a high heritability.
2. A single gene causes narcolepsy in dogs and humans.
3. A family with a very unusual sleep-wake cycle revealed a “clock” gene in humans.

8.4 Intelligence

Intelligence is a vastly complex and variable trait that is subject to many genetic and environmental influences, and also to great subjectivity. Sir Francis Galton, a half first cousin of Charles Darwin, investigated genius, which he defined as “a man endowed with superior faculties.” He identified successful and prominent people in Victorian-era English society, and then assessed success among their relatives. In his 1869 book, *Hereditary Genius*, Galton wrote that relatives of eminent people were more likely to also be successful than people in the general population. The closer the blood relationship, he concluded, the more likely the person was to succeed. This, he claimed, established a hereditary basis for intelligence.

Definitions of intelligence vary. In general, intelligence refers to the ability to reason, learn, remember, connect ideas, deduce, and create. The first intelligence tests, developed in the late nineteenth century, assessed sensory perception and reaction times to various stimuli. In 1904, Alfred Binet at the Sorbonne in France developed a test with verbal, numerical, and pictorial questions, used to predict the success of developmentally disabled youngsters in school. The test was modified at Stanford University to assess white, middle-class Americans. An average score on this “intelligence quotient,” or IQ test, is 100, with two-thirds of all people scoring between 85 and 115 in a bell curve or normal distribution (figure 8.5). An IQ between 50 and 70 is considered mild mental retardation, and below 50, severe mental retardation. In the United States, 3 in 100 individuals have mental retardation. Causes include single-gene and chromosomal disorders, problems during pregnancy or birth, and infection, poisoning, and malnutrition.

IQ has been a fairly accurate predictor of success in school and work. However, low IQ also correlates with many societal situations, such as poverty, a high divorce rate, failure to complete high school, incarceration (males), and having a child out of wedlock (females). Opportunity has a lot to do with intellectual development.

The IQ test consists of short exams that measure verbal fluency, mathematical reasoning, memory, and spatial visualization skills. Because people tend to earn similar scores in all these areas,

psychologists use a general or global intelligence ability, called “g,” to represent the four basic skills that IQ encompasses. In contrast, analysis of personality reveals five contributing factors. The g value is the part of IQ that accounts for differences between individuals based on a generalized intelligence, rather than on enhanced opportunities such as attending classes to boost test-taking skills.

Environment does not seem to play too great a role in IQ differences. Evidence includes the observation that IQ scores of adoptees, with time, become closer to those of their biological parents than to those of their adoptive parents. Heritability studies also reveal a declining environmental impact with age (table 8.2). This makes sense. As a person ages, he or she has more control over the environment, so genetic contributions to intelligence become more prominent.

Researchers have long recognized a genetic explanation for intelligence differences because nearly all syndromes that result from abnormal chromosomes include some degree of mental retardation. The search for single genes that contribute to intelligence differences focuses on proteins that control neurotransmission. For example, a certain variant of a gene encoding neural cellular adhesion molecule (N-CAM) correlates strongly with high IQ. Perhaps this gene variant eases certain neural connections that enhance learning ability. A section of chromosome 4 harbors intelligence-related genes. Genome-wide association studies have identified regions of several chromosomes that likely include genes that affect this hard-to-assess trait.

Age Group	Heritability
Preschoolers	0.4
Adolescents	0.6
Adults	0.8

Key Concepts

1. Intelligence is the use of mental skills to complete complex tasks or solve problems.
2. IQ assesses verbal fluency, mathematical reasoning, memory, and spatial visualization ability.
3. The “g” value measures a general intelligence factor that represents the inherited portion of IQ.
4. Environment has less of an influence on intelligence as a person ages. Individual genes affect intelligence.

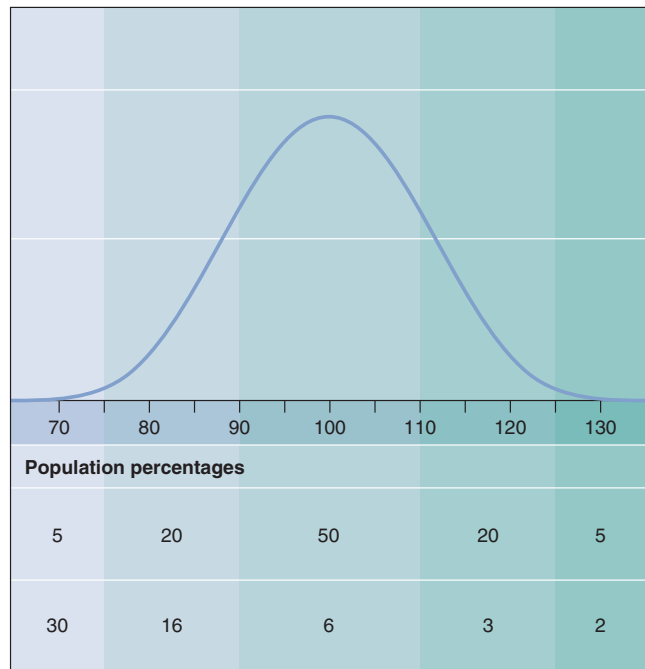


Figure 8.5 Success and IQ. IQ scores predict success in school and the workplace in U.S. society. The bell curve for IQ indicates that most people fall between 85 and 115. However, those living in poverty tend to have lower IQs.

8.5 Drug Addiction

One person sees what a loved one goes through in battling lung cancer and never picks up another cigarette. Another person actually has lung cancer, yet refuses to stop smoking, taking breaks from using her oxygen tank so that she can light up. Evidence is mounting that genes play a large role in making some individuals prone to addiction, and others not.

The Definition of Addiction

Drug addiction is compulsively seeking and taking a drug despite knowing its adverse effects. The two identifying characteristics are tolerance and dependence. Tolerance is the need to take more of the drug to achieve the same effects as time goes on. Dependence is the onset of withdrawal symptoms upon stopping the use of the drug. Both tolerance and dependence contribute to the biological and psychological components of craving the drug. The behavior associated with drug addiction can be extremely difficult to break.

Drug addiction produces long-lasting brain changes. Craving and high risk of relapse remain even after a person has abstained for years. Heritability is 0.4 to 0.6, with a two- to threefold increase in risk among adopted individuals who have one affected biological parent. Twin studies also indicate an inherited component to drug addiction.

Brain imaging techniques localize the “seat” of drug addiction in the brain by highlighting the cell surface receptors

that bind neurotransmitters when a person craves the drug. The brain changes that contribute to addiction occur in parts called the nucleus accumbens, the prefrontal cortex, and the ventral tegmental area, which are part of a larger set of brain structures called the limbic system (**figure 8.6**). The effects of cocaine seem to be largely confined to the nucleus accumbens, whereas alcohol affects the prefrontal cortex.

Although the specific genes and proteins that are implicated in addiction to different substances may vary, several general routes of interference in brain function are at play. Proteins involved in drug addiction are those that

- are part of the production lines for neurotransmitters, such as enzymes;
- form reuptake transporters, which remove excess neurotransmitter from the synapse;
- form receptors on the postsynaptic neuron that are activated or inactivated when specific neurotransmitters bind;
- are part of the signal transduction pathway in the postsynaptic neuron.

Drugs of Abuse

Our ancient ancestors must have discovered that ingesting certain natural substances, particularly from plants, provided a feeling of well-being. That tendency persists today. The degree to which a particular drug is addictive has nothing to do with how a society controls access to it.

Abused drugs are often plant-derived chemicals, such as cocaine, opium, and tetrahydrocannabinol (THC), the main active ingredient in marijuana. These substances bind to receptors on human neurons, which indicates that our bodies have versions of these substances. The human equivalents of the opiates are the endorphins and enkephalins, and the equivalent of THC is anandamide. The endorphins and enkephalins relieve pain. Anandamide modulates how brain cells respond to stimulation by binding to neurotransmitter receptors on

presynaptic (sending) neurons. In contrast, neurotransmitters bind to receptors on postsynaptic neurons.

Amphetamines and LSD produce their effects by binding to receptors on neurons that normally bind neurotransmitters called trace amines. Trace amines are found throughout the brain at low levels, compared to the more abundant neurotransmitters such as dopamine and serotonin. LSD causes effects similar to the symptoms of schizophrenia (see section 8.7), suggesting that the trace amine receptors, which are proteins, may be implicated in the illness.

Candidate gene studies as well as genome-wide association studies are revealing gene variants that people addicted to various drugs share. These inherited factors must be paired with environmental stimuli for addiction to occur. For example, people who are homozygous for the *A1* allele of the dopamine D(2) receptor gene variant are overrepresented among people with alcoholism and people with other addictions. Genome-wide association studies have found 51 chromosomal regions that may include genes that contribute to craving. Finally, studies of gene expression flesh out this picture by providing a real-time view of gene expression changes that happen when a person craves a drug, and then takes it.

Discovering the genetic underpinnings of nicotine addiction is increasing our knowledge of addiction in general, and may have practical consequences. Two facts are clear:

1. The nicotine in tobacco products causes lung cancer.
2. Nicotine is highly addictive.

Genetics explains both.

Each year, 35 million people try to quit smoking, yet only 7 percent succeed. It is easy to see on a whole-body level how this occurs: Nicotine levels peak 10 seconds after an inhalation and the resulting pleasurable release of dopamine in brain cells fades away in minutes. To keep the feeling that researchers call the “reward,” smoking must continue. And it does.

On a molecular level, the tendency to be harmed by nicotine can be traced to a five-part molecular assembly called a nicotinic receptor. The receptor normally binds the neurotransmitter acetylcholine, but it also binds the similarly-shaped nicotine molecule. Certain versions of the receptor bind nicotine very strongly, which triggers a nerve impulse that, in turn, stimulates the pleasurable dopamine release. That may explain the addiction. These receptors are also located on several types of lung cells, where they bind carcinogens. So the nicotine in tobacco causes addiction and susceptibility to lung cancer and it delivers the carcinogens.

Genetics enters the picture in several ways. A different gene encodes each of the five parts of the nicotinic receptor. If two of the five parts are certain variants, then a person experiences desire to continue

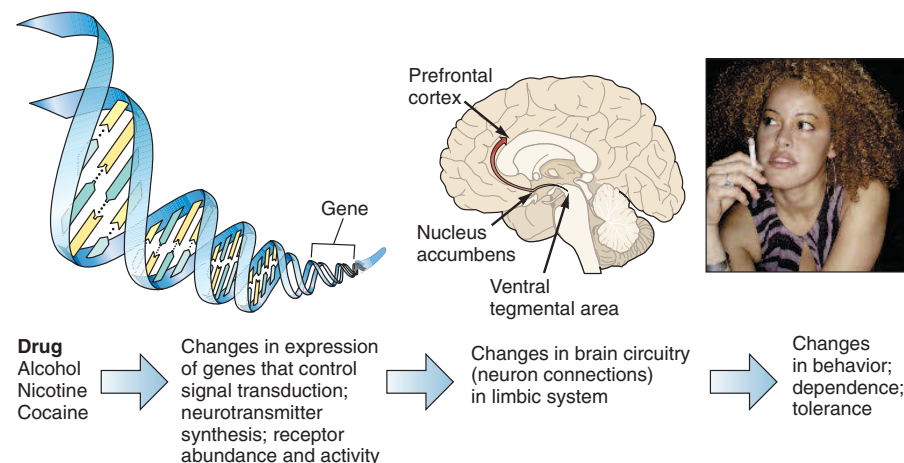


Figure 8.6 The events of addiction. Addiction is manifest at several levels: at the molecular level, in neuron-neuron interactions in the brain, and in behavioral responses.

smoking after the first cigarette, addiction, and increased lung cancer risk. These susceptibility alleles are more common in some populations than others—in one study they were present in 37 percent of Europeans but in none of the Africans. Another gene encodes a protein, called neurexin-1, that ferries the nicotinic receptors to the neuron's surface. Genes also control dopamine synthesis and transport, and how readily the carcinogens cause cancer. One gene variant, for example, greatly increases risk of developing lung cancer—but only if the person smokes.

Key Concepts

1. Drug addiction is dependency on a drug despite knowing the activity is harmful.
2. Structures in the limbic system are directly involved in drug addiction.
3. A candidate gene for addiction encodes the dopamine D(2) receptor.
4. Nicotine binds a receptor that normally binds acetylcholine, causing dopamine release and pleasure.

8.6 Mood Disorders

We all have moods, but mood disorders, which affect millions of people, impair the ability to function on a day-to-day basis. Context is important in evaluating extreme moods. For example, the same symptoms that may lead to a diagnosis of depression are normal in the context of experiencing profound grief. The two most prevalent mood disorders are **major depressive disorder** and **bipolar disorder** (also called manic-depression).

Major Depressive Disorder

Sheeva had trouble remembering how her depression had started. She'd always loved her job at the bookstore after classes at the community college, but it began to annoy her, just as her

classes began to bore her. She had trouble falling and staying asleep, which made her more tired and withdrawn during the day. A few times she'd started to cry while stuck in traffic, not sure why. Then she started having to drag herself out of bed in the morning, completely unmotivated to do anything. She often forgot to eat—food was no longer enjoyable. When she asked her friend Juanita to make some routine phone calls for her, Juanita became alarmed, and gently suggested that Sheeva see a doctor about depression. Sheeva was surprised, but her friend was insistent. After a thorough interview by the physician, Sheeva was indeed diagnosed with major depressive disorder. Within a few weeks of starting treatment with a drug that increases the abundance of the neurotransmitter serotonin in her brain, the fog she had been living with for months began to lift—and she realized she had been sick.

Depression affects 6 percent of the U.S. population at any given time, and affects more women than men. Lifetime risk for the general population is 5 to 10 percent. Often depression is chronic, with acute episodes provoked by stress. It is a serious illness. Fifteen percent of people hospitalized for severe, recurrent depression ultimately end their lives. About half of all people who experience a depressive episode will suffer others. Half of affected individuals do not seek medical help, and among those who do, a third do not respond to drug therapy; those who do may relapse when they discontinue taking an effective drug. Electroconvulsive (shock) therapy can help some patients for whom drugs do not work. For many people, antidepressant treatment is very helpful if paired with psychotherapy.

A likely cause of depression is deficiency of the neurotransmitter serotonin, which affects mood, emotion, appetite, and sleep. Levels of norepinephrine, another type of neurotransmitter, are important as well. In depression, these neurotransmitters become depleted in synapses. Antidepressant drugs called selective serotonin reuptake inhibitors (SSRIs) prevent presynaptic neurons from admitting serotonin from the synapse, leaving more of it available to stimulate the postsynaptic cell (**figure 8.7**). This action apparently offsets the neurotransmitter deficit. Other antidepressants target norepinephrine or

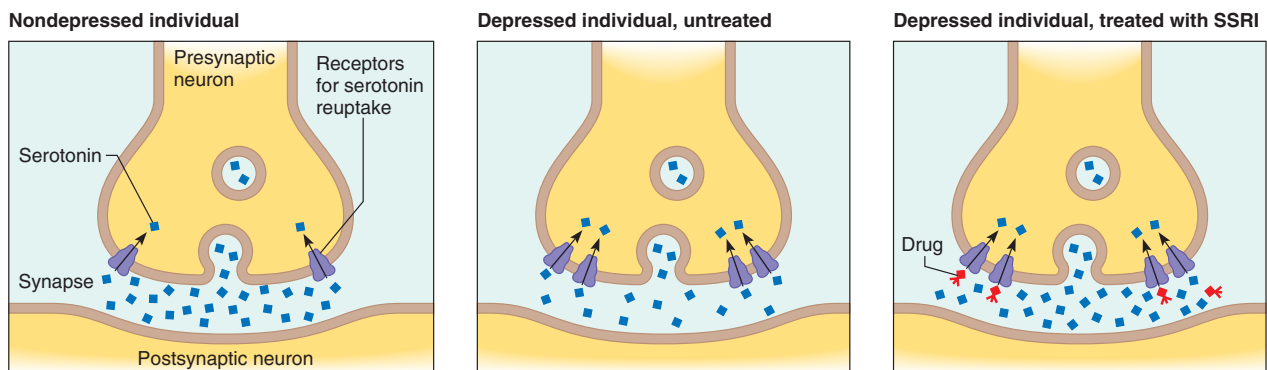


Figure 8.7 How an antidepressant works. Selective serotonin reuptake inhibitors (SSRIs) block the reuptake of serotonin, leaving more of the neurotransmitter in synapses. This corrects a neurotransmitter deficit that presumably causes the symptoms. Overactive or overabundant reuptake receptors can cause the deficit. The precise mechanism of SSRIs is not well understood, and the different drugs may work in slightly different ways.

both serotonin and norepinephrine. SSRIs may begin to produce effects after one week, often enabling a person with moderate or severe depression to return to some activities, but full response can take up to eight weeks.

In the past, physicians would try one antidepressant drug after another, based on experience with other patients, in the hope that one would work. This trial-and-error approach would often take months. Tests of gene expression are being developed that will enable clinicians to choose an antidepressant in a personalized manner, based on which drug is likely to be most effective, with the fewest side effects, in a particular patient.

Bipolar Disorder

Phil Ochs was a political activist in the 1960s who wrote some of the most passionate and compelling anti-war songs of that tumultuous era. Yet his detailed, evocative lyrics were in sharp contrast to his inability to sustain personal relationships. Ochs suffered from bipolar disorder, which made his short life a lonesome highway of extreme ups and downs. His mood first plunged in 1968, in response to the disturbing events of the times—the Democratic National Convention, where he was arrested for protesting, and the assassinations of Martin Luther King Jr. and Robert Kennedy. By 1970, he emerged in full manic mode at Carnegie Hall, dressed in a gold Elvis costume, much to the shock of long-time fans of his folk music (the author was there.) Shortly after this odd performance he declined again, eventually becoming unable to perform, a prisoner to his uncontrollable moods. Ochs committed suicide in 1976.

Bipolar disorder is much rarer than depression, affecting 1 percent of the population and with a general population lifetime risk of 0.5 to 1.0 percent. With this disorder, weeks or months of depression alternate with periods of mania, when the person is hyperactive and restless, and may experience a rush of ideas and excitement. Ideas may be fantastic, and behavior reckless. For example, a person who is normally quiet and frugal might, when manic, suddenly make large monetary donations and spend lavishly—very out-of-character behavior. In one subtype of bipolar disorder, the “up” times are termed hypomania, and they seem more a temporary reprieve from the doldrums than the bizarre behavior of full mania. Bipolar disorder with hypomania may appear to be depression. This is an important distinction because different drugs are used to treat depression and bipolar disorder.

Many gene variants contribute to bipolar disorder. Early genetic studies looked at large Amish families, in whom the manic phase of the disorder was very obvious amid their restrained lifestyle. But studies in different families pointed to different genes behind bipolar disorder. Researchers now realize that the reason why studies disagree is that there are many gene variant combinations that cause or contribute to the phenotype of bipolar disorder, but only a few such variants are seen in any one family. A systematic scrutiny of all studies of the genes behind bipolar disorder suggests as much

as 10 percent of the genome—that is, hundreds of genes—are part of the picture. One researcher compares the approach of considering many studies to a Google search: “The more links there are to a page on the Internet, the more likely it is to come up at the top of your search list. The more experimental lines of evidence for a gene, the higher it comes up in your priority list of genes involved in the disorder.” Question 12 at the chapter’s end lists some of these genes.

Key Concepts

1. Major depressive disorder is more common than bipolar disorder, and is likely caused by deficits of serotonin, norepinephrine, or both.
2. Bipolar disorder is associated with several chromosomal sites, and its genetic roots are difficult to isolate.

8.7 Schizophrenia

Mathematician John Nash Jr. won the Nobel prize in 1994 for game theory based on a short PhD thesis he wrote in 1950, at age 22. He is better known for the portrayal of his paranoid schizophrenia in a book and a film called “A Beautiful Mind.”

Symptoms began in 1959, when Nash’s lectures and conversations would ramble, his thoughts tumbling out so fast, and so disconnected, that people could not understand him. He left academia for the first of several stays in a psychiatric facility. Between hospitalizations, Nash traveled, hearing voices and imagining he was being hunted. Numbers, he thought, carried secret meanings. At various times, Nash had delusions that he was a slave, the messiah, the biblical Job, and the emperor of Antarctica.

John Nash has provided insights into schizophrenia because he got better. In his mid-fifties, he became able to ignore the voices, concentrate, and think mathematically again. Looking back, Nash attributes his professional success to the strange way of thinking that was his schizophrenia. He isn’t alone, fueling the idea that one approach to making schizophrenia easier to live with, for some people, may be to channel the heightened perceptions into art, writing, or music. The list of creative people who had or have schizophrenia includes sir Isaac Newton, Jack Kerouac, Emily Dickinson, and members of the rock bands Fleetwood Mac, Pink Floyd, the Beach Boys, and the Jefferson Airplane.

Schizophrenia is a debilitating loss of the ability to organize thoughts and perceptions, which leads to a withdrawal from reality. Various forms of the condition together affect 1 percent of the world’s population. Ten percent of affected individuals commit suicide.

Identifying genetic contributions to schizophrenia illustrates the difficulties in analyzing a behavioral condition. Some of the symptoms are also associated with other illnesses; many genes cause or contribute to schizophrenia; and several environmental factors mimic the condition.

Signs and Symptoms

The first signs of schizophrenia often affect thinking. In late childhood or early adolescence, a person might develop trouble paying attention in school, and learning may become difficult as memory falters and information-processing skills lag. Symptoms of psychosis begin between ages 17 and 27 for males and 20 and 37 for females, including delusions and hallucinations—sometimes heard, sometimes seen. A person with schizophrenia may hear a voice giving instructions. What others perceive as irrational fears, such as being followed by monsters, are very real to the person with schizophrenia. Meanwhile, cognitive skills continue to decline. Speech reflects the garbled thought process; the person skips from topic to topic with no obvious thread of logic, or displays inappropriate emotional responses, such as laughing at sad news. Artwork by a person with schizophrenia can display the characteristic fragmentation of the mind (**figure 8.8**). (Schizophrenia means “split mind,” but it does not cause a split or multiple personality.)

The course of schizophrenia often plateaus (evens out) or becomes episodic. It is not a continuous decline, as is the case for dementia. Schizophrenia may be misdiagnosed as depression or bipolar disorder, and in fact some of the same gene variants can underlie these conditions. However, schizophrenia primarily affects thinking; these other conditions mostly affect mood. It is a very distinctive mental illness.

Genetic Associations

A heritability of 0.8 and empiric risk values indicate a strong role for genes in causing schizophrenia (**table 8.3**). Because most of the symptoms are behavioral, however, it is possible to develop some of them—such as disordered thinking—from living with and imitating people who have schizophrenia.



Figure 8.8 Schizophrenia alters thinking. People with schizophrenia communicate the disarray of their thoughts in characteristically disjointed drawings.

Table 8.3 Empiric Risks for Schizophrenia

Relationship	Risk
MZ twin	48%
DZ twin	17%
Child	13%
Sibling	9%
Parent	6%
Half sibling	6%
Grandchild	5%
Niece/nephew	4%
Aunt/uncle	2%
First cousin	2%
General population	1%

Although concordance is high, a person who has an identical twin with schizophrenia has a 52 percent chance of *not* developing schizophrenia. Therefore, the condition has a significant environmental component, too.

Researchers hypothesize that dozens of genes may interact with an environmental trigger or triggers to cause schizophrenia (**Reading 8.1**). Genome-wide screens of families with schizophrenia reveal at least twenty-four sites where affected siblings share alleles much more often than the 50 percent of the time that Mendel’s first law predicts.

Environmental Influences

Several environmental factors may increase the risk of developing schizophrenia. One powerful candidate is infection during pregnancy. When a pregnant woman is infected, her immune system bathes the brain of the embryo or fetus with cytokines (molecules that function in signal transduction) that subtly alter brain development.

The idea that maternal infection can sow the seeds for schizophrenia came from observations that an unusually high percentage of people with schizophrenia were born in the winter, especially in the years of flu pandemics. Prenatal exposure to flu virus or the mother’s immune response could contribute to increased risk for development of schizophrenia. The influenza virus can cross the placenta and alter brain cells. Herpes infection at the time of birth is also associated with increased risk of schizophrenia, as is maternal malnutrition, birth complication, fetal oxygen deprivation, and traumatic brain injury to the mother. The opening essay for chapter 7 presents yet another view of schizophrenia—that it is at the opposite end of a continuum from autism, the topic of the next section.

Reading 8.1

Scrutinizing Schizophrenia

One of the difficulties in identifying genes that contribute to behavioral disorders is that these conditions have complex genetic and environmental influences. Both classical family linkage studies and genome-wide association studies are leading researchers to candidate genes for these common conditions. Schizophrenia research offers a good example of how these two approaches both complement and converge: Family studies reveal rare genetic components, whereas genome-wide association studies reveal more common ones. Using results of both types of studies, researchers can “connect the dots” to understand how a disease begins—and this knowledge can suggest new ways to treat the disease.

Family Linkage Studies

At Johns Hopkins University, Akira Sawa’s lab was working with a gene called “disrupted in schizophrenia” (*DISC1*, MIM 605210). Researchers had followed a chromosomal abnormality to track the mutant gene in a large Scottish family in which several members had schizophrenia. *DISC1* protein normally binds a protein called pericentriolar material 1 (*PCM1*, MIM 600299), which is found at the centrosome, a structure that controls how and where microtubules form. The centrosome controls the formation of dendrites, the receiving ends of neurons, and therefore helps to determine how these cells interact.

Elsewhere on campus, Nicholas Katsanis was working on what he thought was an unrelated gene, *BBS4* (MIM 600374). When mutant, it causes Bardet-Biedl syndrome (MIM 209900). This condition affects the eyes, kidneys, and brain. Katsanis noted that the *BBS4* protein binds *PCM1*. It was a third researcher, psychiatrist Nicola Cascella, who connected the work: *DISC1*, *PCM1*, and *BBS4* proteins all work together in a cell, and when the trio is disturbed, neuron-neuron connections change in a way that sets the stage for schizophrenia, or the behavioral symptoms of Bardet-Biedl syndrome.

The next step was to test their hypothesis with experiments. Removing one of the three proteins in cells growing in culture disrupted the functioning of the other two proteins, showing that the three indeed interact. In another set of experiments, developing mice with low levels of either *DISC1* or *BBS4* protein had low levels of *PCM1*, and too few cells in the part of the brain used for thinking and remembering.

Finally, the researchers looked back to families with schizophrenia. In one family, only the affected relatives had a mutation in *PCM1*—a gene not until then associated with schizophrenia. Researchers can now focus on the establishment of specific neural connections in the developing brain as a cause of at least one type of schizophrenia.

A Genome-Wide Approach

People differ not only in their DNA sequences, but in the numbers of short repeated segments. It is a little like the story told in the letters of a book, compared to the number of times the word “the” is used. The genomes of some people who have schizophrenia have more than the usual number of these copy number variants (CNVs), which serve as markers for genes that could contribute to a trait, much as SNPs do (see table 5.3).

Microarray technology enables researchers to probe genomes for small CNVs, which are too tiny to show up as missing or extra segments of chromosomes. In one such study, researchers at the University of Washington in Seattle, comparing the genomes of 150 people with schizophrenia to those of 268 of their healthy relatives, found novel CNVs (small duplications and deletions of the DNA sequence) in 5 percent of the healthy individuals, but in 15 percent of the people with schizophrenia, and in 20 percent of the people who developed schizophrenia at a very young age. Researchers in Iceland had similar results.

Both research groups have repeated their genome-wide CNV studies in many people, confirming that these CNV variants point to possible candidate genes on chromosomes 1, 2, and 15. Researchers are now looking for genes on these chromosomes whose functions are consistent with the phenotype of schizophrenia.

The two types of studies did not discover the same genes for schizophrenia. Taken together, the results may represent rare gene variants with high penetrance in specific families, as well as more common genetic variants with more modest effects. Both approaches—studying families and populations—are valuable in revealing the roles of genes in behavioral disorders.

Key Concepts

1. Schizophrenia affects thinking and causes delusions and hallucinations, usually beginning in young adulthood.
2. Studies have implicated several candidate genes and chromosomal regions as possible causes.
3. A possible environmental influence may be prenatal exposure to the maternal immune system’s response to influenza.

8.8 Autism

Stephen was a “normal” baby in every way. By a year of age he was cruising along the furniture, babbling, smiling, and even saying a few words. By 18 months of age his vocabulary had reached a hundred or so words—plenty to navigate among his toddling friends in daycare. Then, gradually, his language and social skills began to vanish. He stopped initiating conversations, and wouldn’t play with the others, preferring to sit in a corner holding a large rubber ball and rocking back and forth, singing to himself. By age 2, he only used a few words, and

referred to himself as Stephen, rather than “me” or “I.” He did not make eye contact with people, nor did he appear to understand nonverbal cues, such as facial expressions, although he sometimes communicated by using his hands. The preschool teacher alerted the parents as soon as she suspected autism, and a work-up by a medical team confirmed her suspicions. Stephen was able to stay at preschool, as long as he followed a very rigid routine and was permitted to hold his ball. Special education programs in the public school greatly helped.

The autism spectrum disorders (MIM 290850) range from the classical form that Stephen has, to milder forms, including Asperger syndrome, in which language skills remain. These disorders affect three to six children out of every 1,000, striking about four times as many boys as girls. Onset is typically before age 3. About 25 percent of affected children develop seizures as they grow older. Although 70 percent of people with autism have mental retardation; others may be very intelligent.

Discovering causes of autism spectrum disorders has been extremely challenging, because there are many. Attempts to assign environmental triggers have largely failed, except for certain prenatal exposures, such as to rubella (German measles) or the drug valproate. Scientific evidence does not support a link to the mercury compound once used in vaccines—autism has increased since that ingredient has been removed. A better way to identify environmental risk factors for autism spectrum disorders may be to first discover the different genetic subtypes, so that studies compare individuals with the same underlying problem.

Assigning genetic variants to autism has been challenging too. Although siblings of affected children are at a 15 percent risk of being affected, compared to the less than 1 percent for the general population, there are MZ (identical) twins in whom one has autism and one does not. So far more than 30 susceptibility or causative genes have been identified. In about 10 percent of people with autism, the condition is part of a syndrome, including chromosomal disorders such as fragile X syndrome (*Bioethics: Choices for the Future* chapter 12) and Down syndrome (Reading 12.1) or single-gene disorders such as Rett syndrome (Reading 6.2).

In the past chromosomal abnormalities in people with autism led researchers to specific genes. Today genome-wide association studies are supplementing the older twin and chromosomal studies.

The Autism Genetic Resource Exchange Database is looking at 500,000 SNPs in

about 3,000 affected individuals. Another investigation used homozygosity mapping on 88 families from populations in the Middle East where cousin–cousin marriages are part of the culture. Children with autism in the same extended and inbred families tend to have inherited the same blocks of homozygous DNA sequences from their shared ancestors. Researchers identified six parts of the genome where sequences were deleted (absent) in the relatives with autism, then consulted the human genome sequence to identify candidate genes whose functions could explain the symptoms. Mutations in one of these genes, *NHE9*, was then identified in 100 children with autism from families that did not marry among themselves, indicating that it is likely an autism gene.

Studies on two classes of genes that specify proteins that function at the sites where two neurons meet—the synapse—may finally explain how autism develops. These cell adhesion proteins are called neurexins and neuroligins. They are embedded in the membranes of pre- and postsynaptic neurons that use a neurotransmitter called glutamate that spreads excitation (**figure 8.9**). Mutations in these genes cause some cases of autism. Researchers think that these genes, and related ones that control the organization of outgrowths on dendrites, affect the synapses that naturally form in early childhood in response to experiences. This is in contrast to synapses that form during prenatal (before birth) development. The autism spectrum, therefore, may arise from interference with impaired ability, early in life, to form synapses in key brain areas that are necessary for learning and memory.

Disorders that affect our minds are a priority for genome-wide association studies, because they have been so difficult

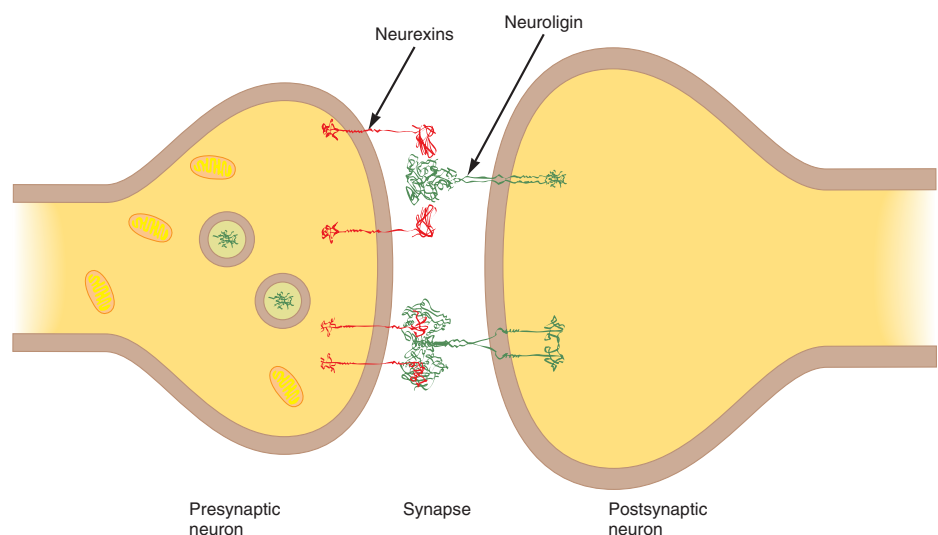


Figure 8.9 Understanding autism. An emerging view of a cause of autism points to changes in the nervous system that occur as a child begins to explore the environment. One such system consists of two types of proteins that strengthen synaptic connections among neurons associated with learning and memory. The proteins are called neurexins (in presynaptic neurons) and neuroligins (in postsynaptic neurons).

to investigate using older methods. A public-private partnership called the Genetic Association Information Network is currently comparing the genomes of 1,000 to 2,000 healthy volunteers with the genomes of equal numbers of volunteers who have any of six disorders chosen because of their impact on public health. Of those six, four are disorders that affect behavior: attention deficit hyperactivity disorder, depression, bipolar disorder, and schizophrenia. The goal is to use clues in genetic differences to develop new prevention and treatment approaches. Stay tuned!

Key Concepts

1. Autism is a loss of language, communication, and social skills beginning in early childhood. Seizures and mental retardation may occur.
2. Many genes contribute to autism risk. Two in particular, which encode the receptor proteins neuroligin and neurexin, may explain how the condition arises from failure of synapses to form that enable a child to integrate experiences.

Summary

8.1 Genes and Behavior

1. Genes contribute to how we respond to environmental stimuli and therefore affect behavior, which includes mood, emotion, intelligence, and personality.
2. Candidate genes for behavioral traits and disorders affect neurotransmission and signal transduction.
3. Analyzing behaviors is difficult because symptoms of different syndromes overlap, study participants can provide biased information, and behaviors can be imitated.
4. Genetic subtypes of behavioral disorders may alter standard psychiatric diagnoses.

8.2 Eating Disorders

5. Eating disorders affect both sexes and are prevalent in the United States and other nations. Twin studies indicate high heritability.
6. Candidate genes for eating disorders include those whose protein products control appetite and the neurotransmitters dopamine and serotonin. Genome-wide association studies point to four genome regions.

8.3 Sleep

7. Twin studies and single-gene disorders that affect the sleep-wake cycle reveal a large inherited component to sleep behavior.
8. A large family with familial advanced sleep phase syndrome enabled researchers to identify the first “clock” gene in humans. The *period* gene enables a person to respond to day and night environmental cues.

8.4 Intelligence

9. Intelligence is difficult to define and measure. The general intelligence (g) value measures the inherited portion of IQ that may underlie population variance in IQ test performance.
10. Heritability for intelligence increases with age, suggesting that environmental factors are more important early in life.
11. Many chromosomal disorders affect intelligence, suggesting high heritability.

8.5 Drug Addiction

12. Drug addiction arises from tolerance and dependence. Addiction produces stable changes in certain parts of the brain.
13. Proteins involved in drug addiction affect neurotransmission and signal transduction.
14. Candidate genes for drug addiction include the dopamine D(2) receptor and variants in nicotinic receptor parts.

8.6 Mood Disorders

15. Major depressive disorder is relatively common and associated with deficits of serotonin and/or norepinephrine.
16. Bipolar disorder is depressive periods and periods of mania or hypomania. Hundreds of genes may raise the risk of developing this disorder. Different families have different combinations of these gene variants, some of which, under certain environmental conditions, can lead to the disorder.

8.7 Schizophrenia

17. Schizophrenia greatly disrupts the ability to think and perceive the world. Onset is typically in early adulthood, and the course is episodic or steady.
18. Empiric risk estimates and heritability indicate a large genetic component.
19. Many genes and environmental influences are associated with schizophrenia.

8.8 Autism

20. Autism is a loss of communication and social skills beginning in early childhood.
21. Neuroligins and neurexins are types of proteins embedded in the cell membranes of certain brain neurons that join across synapses, permitting neural connections to form in response to environmental stimuli. These proteins are abnormal in some cases of autism.

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Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

1. What are the two major types of cells in the brain, and what do they do?
2. Why are behavioral traits nearly always multifactorial?
3. List the pathways or mechanisms that include proteins that, when absent or abnormal, cause variations in behavior.
4. What is the evidence that the Utah family with familial advanced sleep phase syndrome has a genetic condition rather than them all just becoming used to keeping weird hours?
5. Choose a behavior discussed in the chapter and identify the region of the brain and/or a specific molecule that is affected.
6. Why is identifying a candidate gene only a first step in understanding how behavior arises and varies among individuals?
7. Describe three factors that can complicate the investigation of a behavioral trait.
8. Why does the heritability of intelligence decline with age? What were some of the prejudices that were part of studying the inheritance of intelligence?
9. What are the two defining characteristics of drug addiction?
10. Select a drug mentioned in the chapter and explain what it does to the nervous system.
11. Explain how an SSRI antidepressant drug is thought to work.
12. Distinguish between the symptoms of bipolar disorder and schizophrenia.
13. What is the evidence that our bodies have their own uses for cocaine, THC, and opium?
14. What is an environmental factor that may influence the development of schizophrenia?
15. Describe how cell surface receptors are implicated in one of the conditions discussed in the chapter.
16. Select a behavioral disorder mentioned in this text, or another. Compare and contrast the type of information derived from classical studies on families versus genome-wide association studies.

Applied Questions

1. Abnormal serotonin levels contribute to or cause eating disorders, major depressive disorder, and bipolar disorder.
 - a. How can an abnormality in one type of neurotransmitter contribute to different disorders?
 - b. What other neurotransmitter is involved in more than one behavioral disorder?
2. Many older individuals experience advanced sleep phase syndrome. Even though this condition is probably a normal part of aging, how might research on the Utah family with an inherited form of the condition help researchers develop a drug to help the elderly sleep through the night and awaken later in the morning?
3. What nongenetic factor might account for the overrepresentation of minority groups among people with low IQ scores in the United States?
4. How does the subunit structure of the nicotinic receptor provide a mechanism for epistasis? (Epistasis is one gene affecting expression of another; see chapter 5.)
5. A television and film star went into rehab for "sex addiction," much to the embarrassment of his wife. Describe how you would evaluate whether this diagnosis is valid, either in an individual or in a large study.
6. What might be the advantages and disadvantages of a SNP profile done at birth that indicates whether a person is at high risk for developing a drug addiction?
7. In some studies of depression and bipolar disorder, correlations to specific gene variants are only evident when participants are considered in subgroups based on symptoms. What might be a biological basis for this finding?
8. A study found that the risk of schizophrenia among spouses of people with schizophrenia who have no affected blood relatives is 2 percent. What might this indicate about the causes of schizophrenia?
9. Studies indicate that in the United States, the incidence of autism has dramatically increased since 1990.
 - a. Does this finding better support a genetic cause or an environmental cause for autism?
 - b. What is a nongenetic factor that might explain the increased incidence of autism?

10. A “markers for addiction” gene panel scans people’s DNA for variants in the following genes that are associated with addiction tendency. The eleven genes encode proteins that are ion channels, cell surface molecules, receptors, enzymes, or cell adhesion molecules. Look up one of them in *Mendelian*

Inheritance in Man (or elsewhere) and explain its role in increasing the risk of nicotine addiction. The markers are: *CHRNA3*, *CHRNA5*, *CHRN3*, *CLCA1*, *CTNNA3*, *GABRA4*, *KCNJ6*, *NRXN1*, *PIPSK2A*, *TRPC7*, and *VPS13A*.

Web Activities

11. Consult the *Diagnostic and Statistical Manual of Mental Disorders* (DSM-IV). Follow links and list three disorders for which candidate genes have been identified. Discuss how those genes might cause the phenotypes.
12. Researchers looked at the results of all studies searching for the genes behind bipolar disorder. They discovered the following genes, some already known to contribute to the disorder, and some novel:
- Already implicated: *Bdnf*, *Arntl*, *Gsk3b*, *DISC1*, *Nrg1*, *Htr2a*
Novel: *Klf12*, *Aldh1a1*, *A2bp1*, *Ak3l1*, *Rorb*, *Rora*
- a. Look up one gene from each group in MIM and hypothesize how it might contribute to the symptoms of schizophrenia if disrupted.
- b. Psychiatrists consult the DSM to diagnose mental illness, treating such disorders as depression, bipolar disorder, autism, and schizophrenia as distinct and separate. Geneticists look for subtypes of these categories and for distinctive features that might allow better discrimination of the phenotype. Does the fact that the gene *DISC1* seems to be involved in causing both schizophrenia and bipolar disorder argue more for the psychiatrist’s approach, or the geneticist’s approach?
- c. Do you think that a genetic test based on genes involved in bipolar disorder would be helpful? What further information do you think should be included if such data are to be used in a test?
13. Researchers working for an online dating organization analyze information on thousands of individuals for gene variants that affect behaviors, such as empathy, risk-taking, excitability, and loyalty, and categorize the people into four groups. The dating company matches people based on these profiles.
- a. What information would you like to see to determine whether this approach is valid or not for individuals?
- b. What is a benefit of this approach?
- c. What is a limitation or danger of the approach?
14. Two genome-wide association studies sought genes that contribute to intelligence. The first considered 1,000 Mexican-American participants in the San Antonio Family Heart Study, and found evidence pointing to genes on chromosomes 3, 6, and 18. The second study looked at 1,094 children with attention deficit hyperactivity disorder and their siblings, and found gene candidates for intelligence on chromosomes 7, 9, and 14. Suggest two possible reasons why the two studies had different results.
15. Researchers used a version of the placebo effect (being given a sugar pill and thinking it is a drug) on twenty-five people who have social anxiety disorder. The participants had to give a speech at the start and at the end of an 8-week period during which they took a pill that they thought was a medication for their anxiety. Ten of the people improved dramatically, as measured by test scores that rated their anxiety, and brain scans confirmed falling activity in the amygdala, the “fear” area. Next the participants were tested for variants of the gene that encodes tryptophan hydroxylase-2, required to synthesize the mood-related neurotransmitter serotonin. Eight of the ten responders had two copies of a gene variant previously associated with calmness. None of the people who did not respond to the placebo, remaining petrified to speak publicly, had the calm version of the gene.
- a. What are the limitations of this study?
- b. Describe a genome-wide approach to building on this work.
- c. Suggest a practical application of this finding.

Case Studies and Research Results

16. Until the 1990s, bipolar disorder was thought not to affect children under age 18. Psychiatrists today maintain that fewer than 1 percent of children have bipolar disorder. However, the percentage of children being diagnosed with bipolar disorder has soared since 2000, along with the publication of many books written by parents of affected children, and appearances on TV talk shows of affected children and their parents.
- a. Does this pattern of increasing disease incidence suggest a genetic or an environmental cause?
- b. Suggest another explanation for the recent apparent increase in incidence of bipolar disorder in children.
- c. Most children are diagnosed with bipolar disorder based on their answers to questions. What might a genetic diagnosis entail?
17. On the island of Fiji, women once valued having a full figure. Then, in 1995, television arrived, and with it, the show “Melrose Place,” depicting skinny women as the ideal. Within three years, the incidence of eating disorders doubled, with a frightening percentage of the female population regularly vomiting on purpose so that they could continue to eat. Does this information argue more for a genetic or nongenetic cause of eating disorders? How could both influences contribute?

Forensics Focus

18. When 43-year-old F.F. discovered his soon-to-be-ex wife in bed with her boyfriend, he shot and killed them both. The defense ordered a pre-trial forensic psychiatric work-up that included genotyping for the enzyme monoamine oxidase A (*MAOA*, MIM 309850), which breaks down the neurotransmitters serotonin, dopamine, and noradrenalin, and for the serotonin transporter (*SLC6A4*, MIM 182138). A “high-activity” allele for *MAOA* is associated with violence in people who also suffered child abuse. Inheriting one or two “short” alleles of *SLC6A4* is associated with depression and suicidal thoughts, in people who have experienced great stress. F.F. had the high-activity *MAOA* genotype, two short *SLC6A4* alleles, and a lifetime of abuse and stress. However,

the judge ruled that the science was not far enough along to admit the genotyping results as evidence.

- a. Under what conditions or situations do you think it is valid to include genotyping results in cases like this?
- b. In a Dutch family, a mutation disables *MAOA*, causing “a syndrome of borderline mental retardation and abnormal behavior,” according to one report. Family members had committed arson, attempted rape, and shown exhibitionism. How can both high and low levels of an enzyme each cause behavioral problems?
- c. What is a limitation to use of behavioral genotyping in a criminal trial?

PART 3 DNA and Chromosomes



DNA is the genetic material. DNA bursts from this treated bacterial cell. The DNA in a human cell would unravel to nearly 6 feet, yet fit into a cell 6 millionths of a meter across.

CHAPTER

9

DNA Structure
and Replication

Chapter Contents

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DNA *Is* the Hereditary Molecule

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Replication *Is* Semiconservative

Steps of DNA Replication

On the Meaning of Gene

To a biologist, *gene* has a specific definition—a sequence of DNA that tells a cell how to assemble amino acids into a particular protein. To others, “gene” has different meanings:

To folksinger Arlo Guthrie, *gene* means aging without signs of the Huntington disease that claimed his father, legendary folksinger Woody Guthrie.

To rare cats in New England, *gene* means extra toes.

To Adolph Hitler and others who have dehumanized those not like themselves, the concept of *gene* was abused to justify genocide.

To a smoker, a *gene* may mean lung cancer develops.

To a redhead in a family of brunettes, *gene* means an attractive variant.

To a woman whose mother and sisters had breast cancer, *gene* means escape from their fate—and survivor guilt.

To a lucky few, *gene* means a mutation that locks HIV out of their cells.

To people with diabetes, *gene* means safer insulin.

To an elephant on the African savannah and one that lives in the forest, *gene* means that they cannot mate with each other.

To a forensic entomologist, *gene* means a clue to the identity of a criminal in the guts of maggots devouring a corpse.

To scientists-turned-entrepreneurs, *gene* means money.

Collectively, our genes mean that we are very much more alike than different from one another.

9.1 Experiments Identify and Describe the Genetic Material

“A genetic material must carry out two jobs: duplicate itself and control the development of the rest of the cell in a specific way,” wrote Francis Crick, codiscoverer with James Watson of the three-dimensional structure of DNA in 1953. Only DNA can do this.

DNA was first described in the mid-eighteenth century, when Swiss physician and biochemist Friedrich Miescher isolated nuclei from white blood cells in pus on soiled bandages. In the nuclei, he discovered an unusual acidic substance containing nitrogen and phosphorus. He and others found it in cells from a variety of sources. Because the material resided in cell nuclei, Miescher called it *nuclein* in an 1871 paper; subsequently, it was called a nucleic acid. Few people appreciated the importance of Miescher’s discovery at the time, when inherited disease was widely blamed on protein.

In 1902, English physician Archibald Garrod was the first to provide evidence linking inherited disease and protein. He noted that people who had certain inborn errors of metabolism lacked certain enzymes. One of the first inborn errors that he described was alkaptonuria, the subject of the chapter 5 opening essay. Other researchers added evidence of a link between heredity and enzymes from other species, such as fruit flies with unusual eye colors and bread molds with nutritional deficiencies. Both organisms had absent or abnormal specific

enzymes. As researchers wondered what, precisely, was the connection between enzymes and heredity, they returned to Miescher’s discovery of nucleic acids.

DNA Is the Hereditary Molecule

In 1928, English microbiologist Frederick Griffith took the first step in identifying DNA as the genetic material. Griffith noticed that mice with a certain form of pneumonia harbored one of two types of *Diplococcus pneumoniae* bacteria. Type R bacteria were rough in texture. Type S bacteria were smooth because they are enclosed in a polysaccharide (a type of carbohydrate) capsule. Mice injected with type R bacteria did not develop pneumonia, but mice injected with type S did. The polysaccharide coat shielded the bacteria from the mouse immune system, enabling them to cause severe (virulent) infection.

When type S bacteria were heated—which killed them but left their DNA intact—they no longer could cause pneumonia in mice. However, when Griffith injected mice with a mixture of type R bacteria plus heat-killed type S bacteria—neither of which, alone, was deadly to the mice—the mice died of pneumonia (**figure 9.1**). Their bodies contained live type S bacteria, encased in polysaccharide. Griffith termed the apparent conversion of one bacterial type into another “transformation.” How did it happen? What component of the dead, smooth bacteria transformed type R to type S?

U.S. physicians Oswald Avery, Colin MacLeod, and Maclyn McCarty hypothesized that a nucleic acid might be

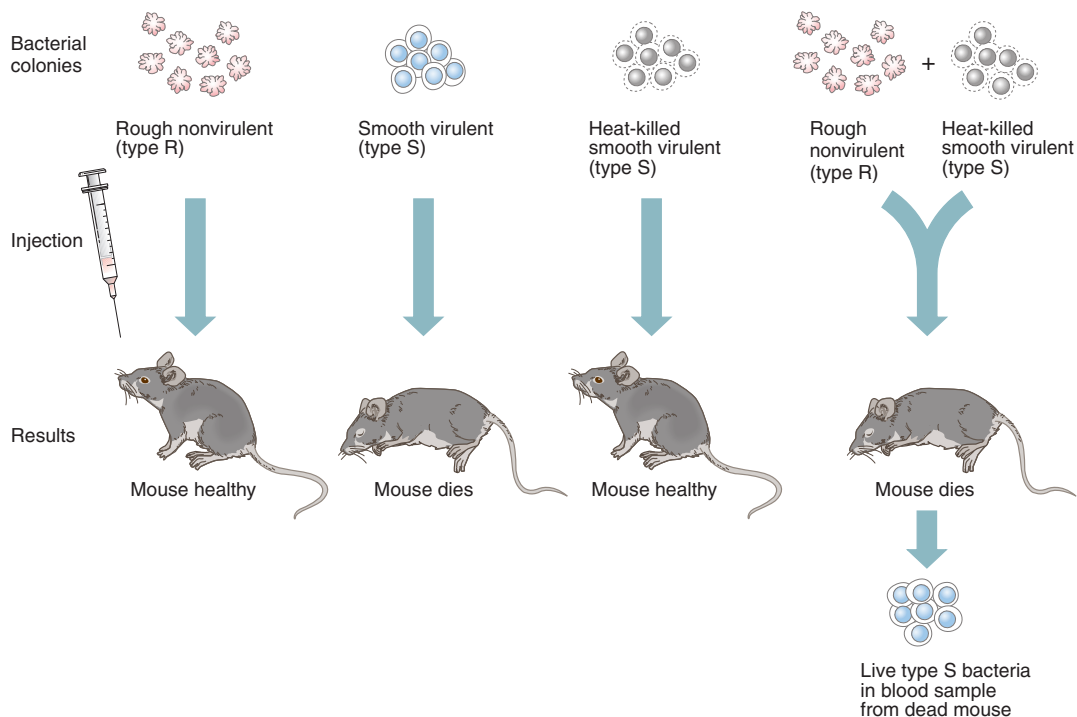


Figure 9.1 Discovery of bacterial transformation. Griffith’s experiments showed that a molecule in a lethal type of bacteria can transform nonkilling (nonvirulent) bacteria into killers (virulent).

the “transforming principle.” They observed that treating broken-open type S bacteria with a protease—an enzyme that dismantles protein—did not prevent the transformation of a nonvirulent to a virulent strain, but treating it with deoxyribonuclease (or DNase), an enzyme that dismantles DNA only, did disrupt transformation. In 1944, they confirmed that DNA transformed the bacteria. They isolated DNA from heat-killed type S bacteria and injected it with type R bacteria into mice (**figure 9.2**). The mice died, and their bodies contained active type S bacteria. The conclusion: DNA passed from type S bacteria into type R, enabling the type R to manufacture the smooth coat necessary for infection.

Protein Is Not the Hereditary Molecule

Science seeks answers by eliminating explanations. To identify the genetic material, researchers also had to show that protein does *not* transmit genetic information. To do this, in 1953, U.S. microbiologists Alfred Hershey and Martha Chase used *E. coli* bacteria infected with a virus that consisted of a protein “head” surrounding DNA. Viruses infect bacterial cells by injecting their DNA (or RNA) into them. Infected bacteria may then produce many more viruses. The viral protein coats remain outside the bacterial cells.

Researchers can analyze viruses by growing them on culture medium that contains a radioactive chemical that the viruses take up. The “labeled” viral nucleic acid then emits radiation, which can be detected in several ways. When Hershey and Chase grew viruses with radioactive sulfur, the viral protein coats emitted radioactivity. When they repeated the experiment with radioactive phosphorus, the viral DNA emitted radioactivity. If protein is the genetic material, then the infected bacteria would have radioactive sulfur. But if DNA is the genetic material, then the bacteria would have radioactive phosphorus.

Hershey and Chase labeled two batches of virus, growing one in a medium containing radioactive sulfur (designated ^{35}S) and the other in a medium containing radioactive phosphorus (designated ^{32}P). The viruses grown on sulfur had their protein

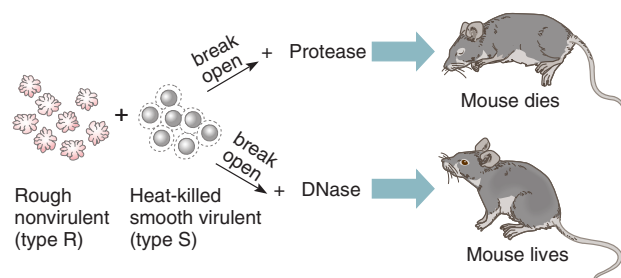


Figure 9.2 DNA is the “transforming principle.” Avery, MacLeod, and McCarty identified DNA as Griffith’s transforming principle. By adding enzymes that either destroy proteins (protease) or DNA (deoxyribonuclease or DNase) to bacteria that were broken apart to release their contents, they demonstrated that DNA transforms bacteria—and that protein does not.

marked, but not their DNA, because protein incorporates sulfur but DNA does not. Conversely, the viruses grown on labeled phosphorus had their DNA marked, but not their protein, because this element is found in DNA but not protein. (Miescher had noted phosphorus in DNA from soiled bandages.)

After allowing several minutes for the virus particles to bind to the bacteria and inject their DNA into them, Hershey and Chase agitated each mixture in a blender, shaking free the empty virus protein coats. The contents of each blender were collected in test tubes, then centrifuged (spun at high speed). This settled the bacteria at the bottom of each tube because the lighter virus coats drift down more slowly than bacteria.

At the end of the procedure, Hershey and Chase examined fractions containing the virus coats from the top of each test tube and the infected bacteria that had settled to the bottom (**figure 9.3**). In the tube containing viruses labeled with sulfur, the virus coats were radioactive, but the virus-infected bacteria, containing viral DNA, were not. In the other tube, where the virus had incorporated radioactive phosphorus, the virus coats carried no radioactive label, but the infected bacteria were radioactive. Therefore, the part of the virus that could enter bacteria and direct them to mass produce more virus was the part that had incorporated phosphorus—the DNA. The genetic material was DNA, and not protein.

Discovering the Structure of DNA

In 1909, Russian-American biochemist Phoebus Levene identified the 5-carbon sugar **ribose** as part of some nucleic acids, and in 1929, he discovered a similar sugar—**deoxyribose**—in other nucleic acids. He had revealed a major chemical distinction between RNA and DNA: RNA has ribose, and DNA has deoxyribose.

Levene then discovered that the three parts of a nucleic acid—a sugar, a nitrogen-containing base, and a phosphorus-containing component—are present in equal proportions. He deduced that a nucleic acid building block must contain one of each component. Furthermore, although the sugar and phosphate portions were always the same, the nitrogen-containing bases were of four types. Scientists at first thought that the bases were present in equal amounts, but if this were so, DNA could not encode as much information as it could if the number of each base type varied. Imagine how much less useful a written language would be if it had to use all the letters with equal frequency.

In the early 1950s, two lines of experimental evidence converged to provide the direct clues that finally revealed DNA’s structure. Austrian-American biochemist Erwin Chargaff showed that DNA in several species contains equal amounts of the bases **adenine** (A) and **thymine** (T) and equal amounts of the bases **guanine** (G) and **cytosine** (C). Next, English physicist Maurice Wilkins and English chemist Rosalind Franklin bombarded DNA with X rays using a technique called X-ray diffraction, then deduced the overall structure of the molecule from the patterns in which the X rays were deflected.

Rosalind Franklin provided a pivotal clue to the structure of DNA to Watson and Crick. She distinguished two forms of

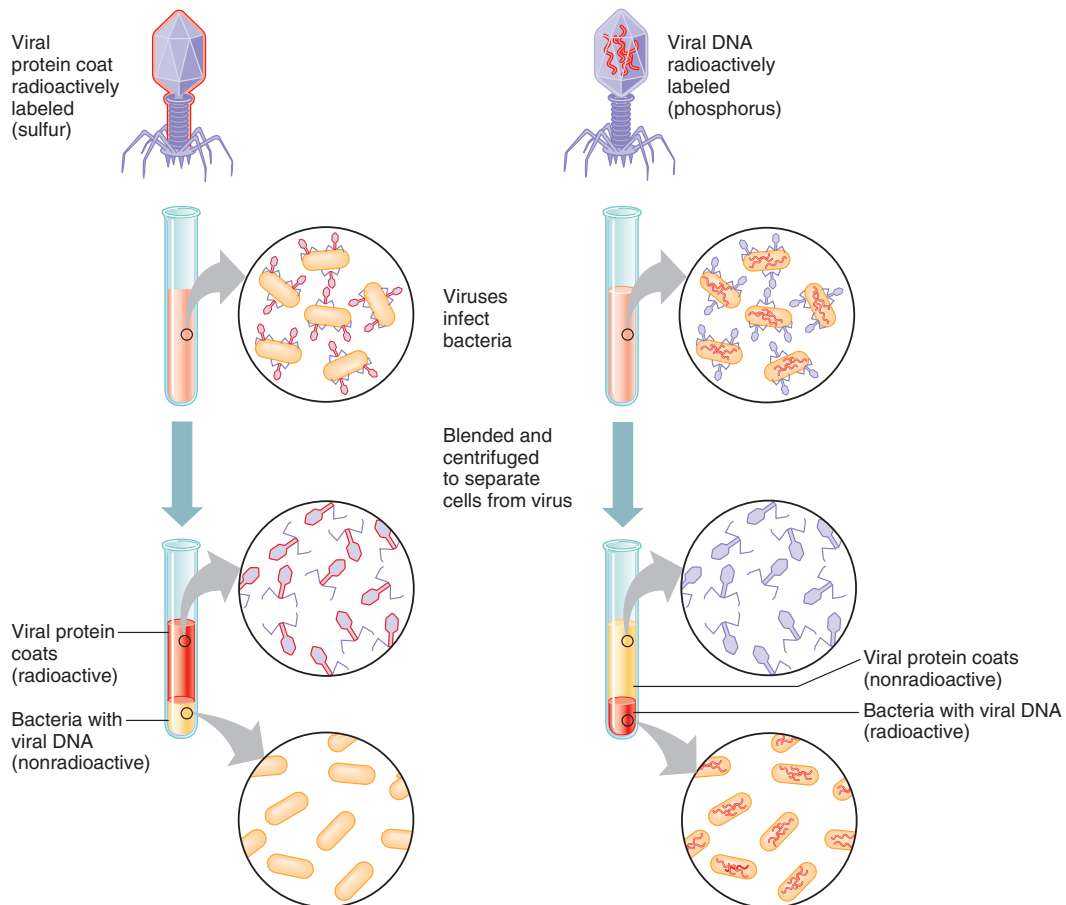
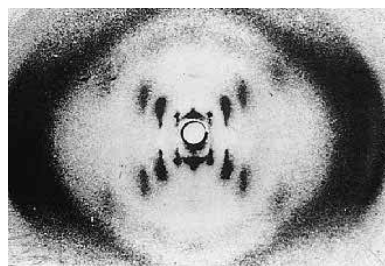


Figure 9.3 DNA is the hereditary material; protein is not. Hershey and Chase used different radioactive molecules to distinguish the viral protein coat from the genetic material (DNA). These “blender experiments” showed that the virus transfers DNA, and not protein, to the bacterium. Therefore, DNA is the genetic material. The blender experiments used particular types of sulfur and phosphorus atoms that emit detectable radiation.

DNA—a dry, crystalline “A” form, which had been well-studied, and the wetter type seen in cells, the “B” form. It took her 100 hours to obtain “photo 51” of the B form in May 1952 (**figure 9.4**). Its remarkable symmetry told Franklin that the molecule was a sleek helix, and revealed the position of the phosphates. She had long thought of DNA as a candidate for the genetic material. A lab notebook from her college days in 1939 bears the comment, “Geometrical basis for inheritance?” next to an illustration of a nucleic acid. By early 1953, she was very close to deducing the entire structure. On January 30, Wilkins showed Franklin’s photo 51 to Watson, who instantly recognized its importance.

The race was on. During February, famed biochemist Linus Pauling suggested a triple helix structure for DNA. Meanwhile, Watson and Crick, certain of the sugar-phosphate



a.

Figure 9.4 Deciphering DNA structure. (a) Rosalind Franklin’s “photo 51” of B DNA was critical to Watson and Crick’s deduction of the three-dimensional structure of the molecule. The “X” in the center indicates a helix, and the darkened regions reveal symmetrically organized subunits. (b) Franklin died very young, of cancer.



b. Rosalind Franklin 1920–1958



Figure 9.5 Watson and Crick. Prints of this famed, if posed, photo fetched a high price when signed and sold at celebrations of DNA's fiftieth anniversary in 2003. Crick, told to point to the model, picked up a slide rule.

backbone largely from photo 51, turned their attention to the bases. Ironically, their eureka moment occurred not with sophisticated chemistry or crystallography, but while working with cardboard cutouts of the DNA components.

On Saturday morning, February 28, Watson arrived early for a meeting with Crick. While he was waiting, he played with cardboard cutouts of the four DNA bases, pairing A with A, then A with G. When he assembled A next to T, and G next to C, he noted the similar shapes, and suddenly all the pieces fit. He had been modeling the chemical attractions between the bases that create the sleek helix. When Crick arrived 40 minutes later, the two quickly realized they had solved the puzzle of DNA's structure (**figure 9.5**).

Watson, Crick, and Wilkins eventually received the Nobel prize. In 1958, Franklin died at the age of 37 from ovarian cancer,

and the Nobel can only be awarded to a living person. In recent years, she has become a heroine for her long-unappreciated role in deciphering the structure of DNA. **Table 9.1** summarizes some of the experiments that led to the discovery.

Key Concepts

1. DNA replicates, and contains information for protein synthesis.
2. Miescher isolated DNA in 1869.
3. Garrod linked heredity to enzymes.
4. In the 1940s, Griffith identified a substance capable of transmitting infectiousness, which Avery, MacLeod, and McCarty showed was DNA.
5. Hershey and Chase confirmed that DNA, and not protein, is the genetic material.
6. Using Chargaff's discovery that the number of As equals the number of Ts, and the number of Gs equals the number of Cs, with Franklin's discovery that DNA is regular and symmetrical, Watson and Crick deciphered the structure of DNA.

9.2 DNA Structure

A **gene** is a section of a DNA molecule whose sequence of building blocks specifies the sequence of amino acids in a particular protein. The activity of the protein imparts the phenotype. The fact that different building blocks combine to form nucleic acids enables them to carry information, as the letters of an alphabet combine to form words. DNA also encodes RNA that does not specify a protein, but instead assists in protein synthesis or controls gene expression. These DNA and RNA sequences are discussed in chapters 10 and 11.

Inherited traits are diverse because proteins have diverse functions (see table 10.1). Malfunctioning or inactive proteins, which reflect genetic defects, can devastate health. Most of the

Table 9.1 The Road to the Double Helix

Investigator	Contribution	Timeline
Friedrich Miescher	Isolated nuclein in white blood cell nuclei	1869
Frederick Griffith	Transferred killing ability between types of bacteria	1928
Oswald Avery, Colin MacLeod, and Maclyn McCarty	Discovered that DNA transmits killing ability in bacteria	1940s
Alfred Hershey and Martha Chase	Determined that the part of a virus that infects and replicates is its nucleic acid and not its protein	1950
Phoebus Levene, Erwin Chargaff, Maurice Wilkins, and Rosalind Franklin	Discovered DNA components, proportions, and positions	1909–early 1950s
James Watson and Francis Crick	Elucidated DNA's three-dimensional structure	1953
James Watson	Had his genome sequenced	2008

amino acids that are assembled into proteins come from the diet or from breaking down proteins in the cell. The body synthesizes the others.

The structure of DNA is easiest to understand if we begin with the smallest components. A single building block of DNA is a **nucleotide**. It consists of one deoxyribose sugar, one phosphate group (a phosphorus atom bonded to four oxygen atoms), and one nitrogenous base. **Figure 9.6** shows the chemical structures of the four types of bases, and **figure 9.7** shows one of them as part of a nucleotide. Adenine (A) and guanine (G) are **purines**, which have a two-ring structure. Cytosine (C) and thymine (T) are **pyrimidines**, which have a single-ring structure.

The bases are the information-containing parts of DNA because they form sequences. DNA sequences are measured in numbers of base pairs. The terms kilobase (kb) and megabase (mb) are used to abbreviate a thousand and a million DNA bases, respectively. A particular gene, for example, may be “1,400 base pairs long.”

Nucleotides join into long chains when chemical bonds form between the deoxyribose sugars and the phosphates. This creates a continuous **sugar-phosphate backbone** (**figure 9.8**). Two such chains of nucleotides align head-to-toe, as **figure 9.9a** depicts. M. C. Escher’s drawing of hands in **figure 9.9b** resembles the spatial relationship of the two strands of the DNA double helix.

The opposing orientation of the two nucleotide chains in a DNA molecule is called **antiparallelism**. It derives from the structure of the sugar-phosphate backbone. Antiparallelism becomes evident when the carbons of the sugars are assigned numbers to indicate their positions in the molecule (**figure 9.10**).

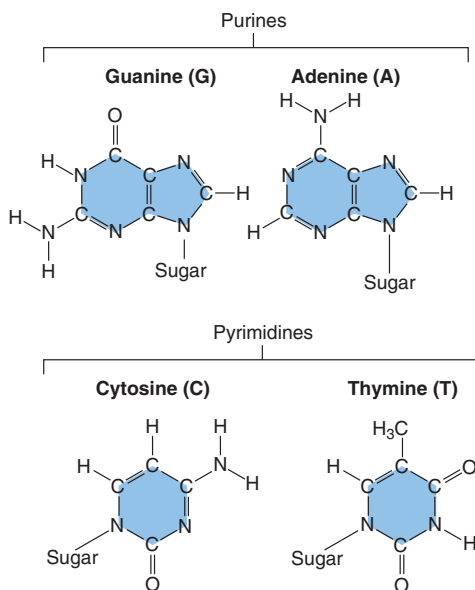


Figure 9.6 DNA bases are the informational parts of nucleotides. Adenine and guanine are purines, each composed of a six-membered organic ring plus a five-membered ring. Cytosine and thymine are pyrimidines, each with a single six-membered ring. (Within the molecules, C, H, N, O, and P are atoms of carbon, hydrogen, nitrogen, oxygen, and phosphorus, respectively.)

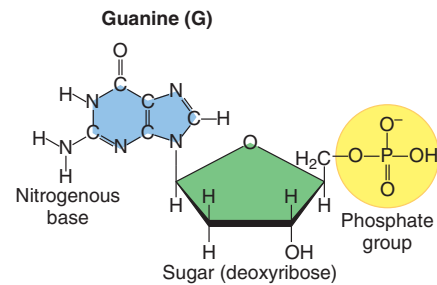


Figure 9.7 Nucleotides. A nucleotide of a nucleic acid consists of a 5-carbon sugar, a phosphate group, and an organic, nitrogenous base (G, A, C, or T).

The carbons are numbered from 1 to 5, starting with the first carbon moving clockwise from the oxygen in each sugar in **figure 9.11**. One chain runs from the #5 carbon (top of the figure) to the #3 carbon, but the chain aligned with it runs from the #3 to the #5 carbon. These ends are depicted as “5′” and “3′”, pronounced “5 prime” and “3 prime.”

The symmetrical DNA double helix forms when nucleotides containing A pair with those containing T, and nucleotides containing G pair with those carrying C. Because purines have two rings and pyrimidines one, the consistent pairing of a purine with a pyrimidine ensures that the double helix has the same width throughout, as Watson discovered using cardboard cutouts. These specific purine-pyrimidine couples are called **complementary base pairs**. Chemical attractions called hydrogen bonds hold the base pairs together. They are weak individually, but over the many bases of a DNA molecule impart great strength. Two hydrogen bonds join A and T, and three hydrogen bonds join G and C, as **figure 9.12**

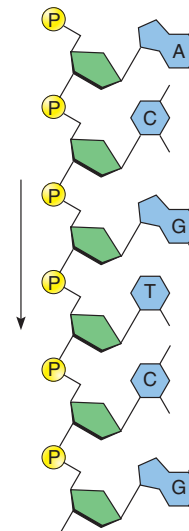
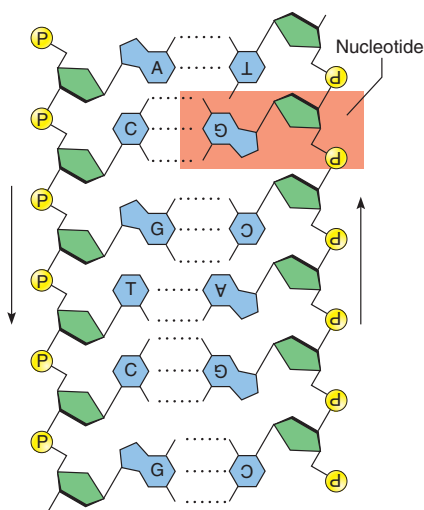
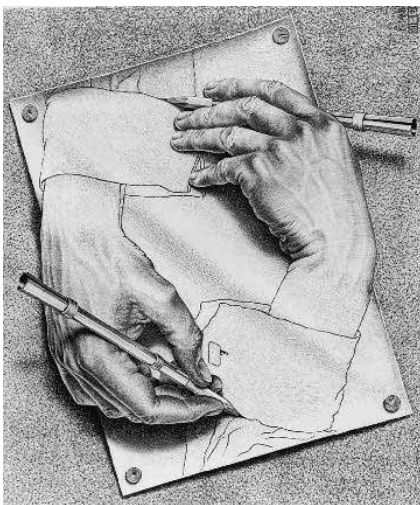


Figure 9.8 A chain of nucleotides. A single DNA strand consists of a chain of nucleotides that forms when the deoxyribose sugars (green) and phosphates (yellow) bond to create a sugar-phosphate backbone. The bases A, C, G, and T are blue.



a.



b.

Figure 9.9 DNA consists of two chains of nucleotides.

(a) Hydrogen bonds hold the nitrogenous bases of one strand to the nitrogenous bases of the second strand (dotted lines). Note that the sugars point in opposite directions—that is, the strands are antiparallel. (b) Artist M. C. Escher captured the essence of antiparallelism in his depiction of hands.

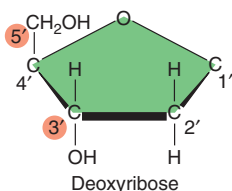


Figure 9.10 Antiparallelism. The antiparallel nature of the DNA double helix becomes apparent when the carbons in the sugar are numbered.

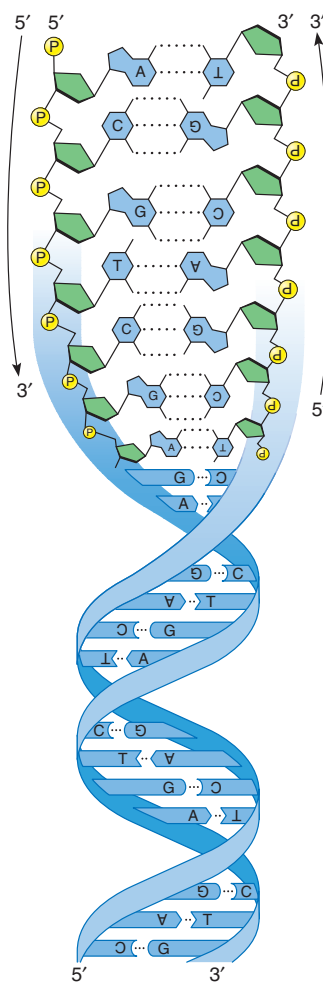


Figure 9.11 DNA is directional. Antiparallelism in a DNA molecule arises from the orientation of the deoxyribose sugars. One-half of the double helix runs in a 5' to 3' direction, and the other half runs in a 3' to 5' direction.

shows. Finally, DNA forms a double helix when the antiparallel, base-paired strands twist about one another in a regular fashion. The double-stranded, helical structure of DNA gives it great strength—50 times the strength of single-stranded DNA, which would not form a helix.

DNA molecules are incredibly long. The DNA of the smallest human chromosome, if stretched out, would be 14 millimeters long. But it is packed into a chromosome that, during cell division, is only 2 micrometers long. This means that the DNA molecule must fold so tightly that its compacted length shrinks by a factor of 7,000:

$$\left(\frac{14 \times 10^{-3} \text{ meters}}{2 \times 10^{-6} \text{ meters}} \right)$$

Various types of proteins compress the DNA without damaging or tangling it. Scaffold proteins form frameworks that guide DNA strands. Then, the DNA coils around proteins called

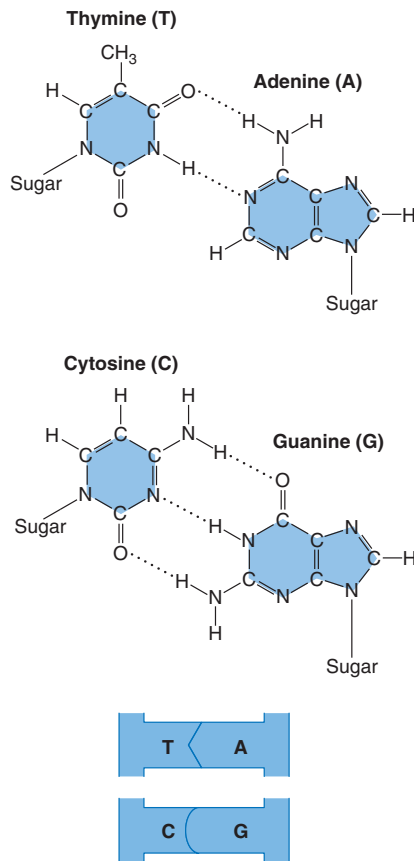


Figure 9.12 DNA base pairs. The key to the constant width of the DNA double helix is the pairing of purines with pyrimidines. Two hydrogen bonds join adenine and thymine; three hydrogen bonds link cytosine and guanine.

histones, forming a beads-on-a-string-like structure. The bead part is called a **nucleosome**. It is a little like wrapping a very long, thin piece of thread around your fingers, to keep it from unraveling and tangling. DNA wraps at several levels, until it is compacted into a chromosome (**figure 9.13**). Specifically, a nucleosome forms around packets of eight histone proteins (a pair of each of four types). A fifth type of histone protein anchors nucleosomes to short “linker” regions of DNA, which then tighten the nucleosomes into fibers 30 nanometers (nm) in diameter. As a result, at any given time, only small sections of the DNA double helix are exposed. Chemical modification of the histones controls when particular DNA sequences are accessible. (This is discussed further in chapter 11.) DNA also unwinds locally when it replicates.

Altogether, the chromosome substance is called **chromatin**, which means “colored material.” Chromatin is not just DNA; it is about 30 percent histone proteins, 30 percent DNA binding proteins, 30 percent DNA, and 10 percent RNA. Specific points along the chromatin attach it, in great loops, to the inner face of the nuclear membrane, placing particular chromosome parts in particular locations in the nucleus.

Key Concepts

1. The DNA double helix’s backbone is alternating deoxyribose and phosphate held together by complementary pairs of A-T and G-C bases. A and G are purines; T and C are pyrimidines.
2. The DNA double helix is antiparallel, its strands running in an opposite head-to-toe manner.
3. DNA winds tightly about histone proteins, forming nucleosomes, which in turn wind tighter, forming chromatin.

9.3 DNA Replication— Maintaining Genetic Information

As soon as Watson and Crick deciphered the structure of DNA, its mechanism for replication became obvious. They ended their report on the structure of DNA with the statement, “*It has not escaped our notice that the specific pairing we have postulated immediately suggests a possible copying mechanism for the genetic material.*”

Thirty years after Watson and Crick’s discovery, biochemist Kary Mullis invented a way to harness DNA replication to mass-produce selected DNA sequences in the laboratory. His polymerase chain reaction (PCR) revolutionized biology research and inspired several other types of DNA amplification technologies. PCR is discussed in section 19.2.

Replication Is Semiconservative

Watson and Crick envisioned the two halves of the DNA double helix unwinding and separating, exposing unpaired bases that would attract their complements. In this way, two double helices would form from one. This route to replication is called **semiconservative**, because each new DNA double helix conserves half of the original. However, separating the long strands posed a huge physical challenge, a little like having to keep two pieces of thread the length of a football field from tangling.

At first, some researchers suggested that DNA might replicate in any of three possible ways:

- semiconservative,
- conservative, with one double helix specifying creation of a second double helix, and
- dispersive, with a double helix shattering into pieces that would join with newly synthesized DNA pieces to form two molecules.

Figure 9.14 shows these three models.

It would take experiments to see which of the three possible ways that DNA can replicate is what really happens. In 1941, English geneticist J. B. S. Haldane had an idea. He wrote: *How can one distinguish between model and copy? Perhaps you could use heavy nitrogen atoms in the food supplied to your cell, hoping that the “copy” genes would contain it while the models did not.*

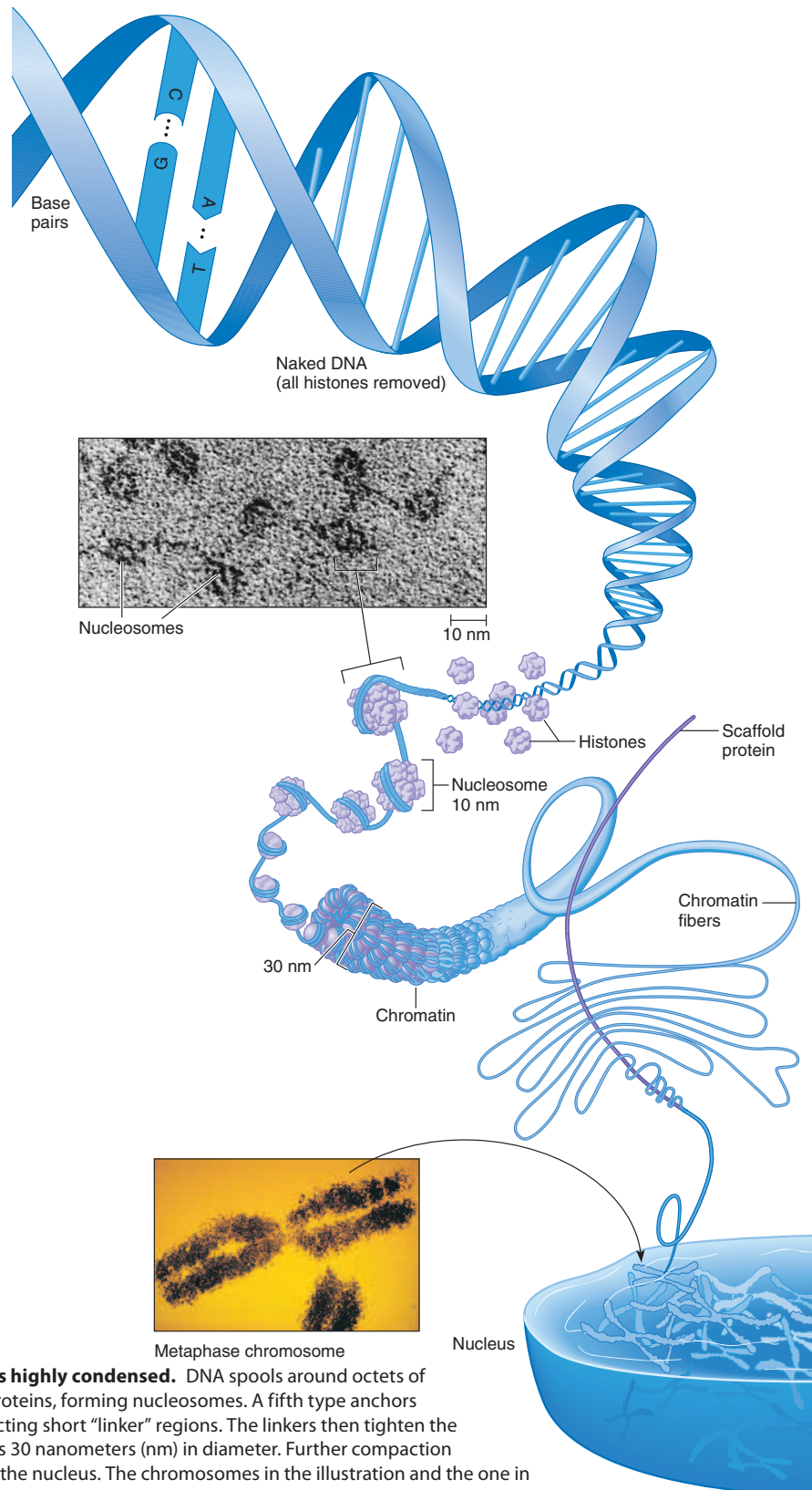


Figure 9.13 DNA is highly condensed. DNA spools around octets of four types of histone proteins, forming nucleosomes. A fifth type anchors nucleosomes to connecting short “linker” regions. The linkers then tighten the nucleosomes into fibers 30 nanometers (nm) in diameter. Further compaction squeezes the DNA into the nucleus. The chromosomes in the illustration and the one in the photograph are in the replicated form. That is, each consists of two DNA molecules, joined at the centromere.

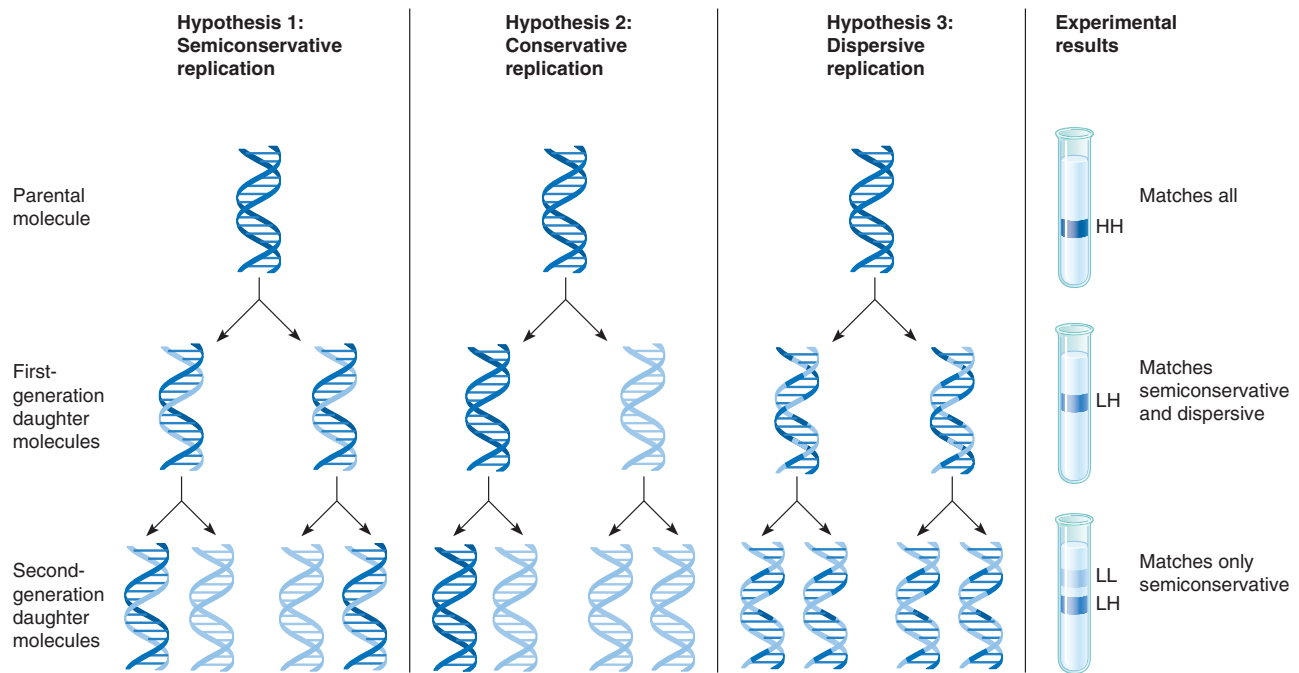


Figure 9.14 Three models for DNA replication. Density shift experiments distinguished the three hypothesized mechanisms of DNA replication. DNA molecules containing light nitrogen are designated “LL” and those with heavy nitrogen, “HH.” Molecules containing both isotopes are designated “LH.” These experiments established that DNA replication is semiconservative. The first three columns illustrate how parental and daughter DNA would be distributed in each of the three mechanisms of DNA replication. The fourth column depicts the density of the DNA at each stage for each of the three hypothesized replication mechanisms.

In 1957, two young researchers, Matthew Meselson and Franklin Stahl, tried Haldane’s experiment using bacteria. Their experimental results not only supported one hypothesis (semiconservative replication), but disproved the other two (conservative replication and dispersive replication).

Meselson and Stahl labeled DNA newly synthesized by bacteria with heavy nitrogen (^{15}N) in the media. The DNA could then be distinguished from older DNA that had been synthesized with the more common lighter form, ^{14}N . The idea was that DNA that incorporated the heavy nitrogen could be separated from newly synthesized DNA that incorporated the normal lighter nitrogen by its greater density. DNA in which one-half of the double helix was light and one-half heavy would be of intermediate density.

In their “density shift” experiments, Meselson and Stahl grew cells on media with heavy nitrogen and then shifted the cells to media with light nitrogen. They traced replicating DNA through several cell divisions. The researchers grew cells, broke them open, extracted DNA, and spun it in a centrifuge. The heavier DNA sank to the bottom of the centrifuge tube, the light DNA rose to the top, and the heavy-light double helices settled in the middle area of the tube.

Meselson and Stahl grew *E. coli* on media containing ^{15}N for several generations, making all of the DNA heavy. They knew this because only “heavy-heavy” molecules appeared in the tube after centrifugation. They then shifted the bacteria to media containing ^{14}N , allowing enough time for the bacteria to divide only once (about 30 minutes).

When Meselson and Stahl collected the DNA after one generation and centrifuged it, the double helices were all of intermediate density. The DNA settled in the middle of the tube, indicating that the molecules contained half ^{14}N and half ^{15}N . This pattern was consistent with either semiconservative DNA replication or a dispersive mechanism. In contrast, the result of conservative replication would have been one band of material in the tube completely labeled with ^{15}N , corresponding to one double helix, and another totally “light” band containing ^{14}N only, corresponding to the other double helix. This did not happen, eliminating conservative replication.

To definitively distinguish among the three routes to DNA replication, Meselson and Stahl extended the experiment one more generation. If the semiconservative mechanism held up, each hybrid (half ^{14}N and half ^{15}N) double helix present after the first generation following the shift to ^{14}N medium would separate and assemble a new half from bases labeled only with ^{14}N . This would produce two double helices with one ^{15}N (heavy) and one ^{14}N (light) chain, plus two double helices containing only ^{14}N . The tube would have one heavy-light band and one light-light band. This is indeed what Meselson and Stahl saw.

The conservative mechanism would have yielded two bands in the tube in the third generation, indicating three completely light double helices for every completely heavy one, as the bottom portion of the hypothesis 2 column indicates in figure 9.14. The third generation for the dispersive model would have been a single large band, somewhat higher than the second-generation band because additional ^{14}N would have been randomly incorporated into the DNA.

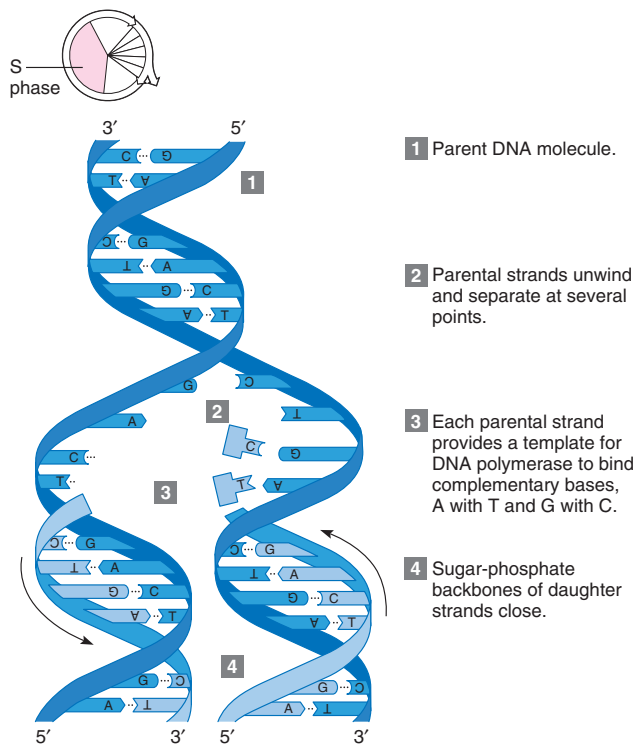


Figure 9.15 Overview of DNA replication.

After experiments demonstrated the semiconservative nature of DNA replication, the next challenge was to decipher the steps of the process.

Steps of DNA Replication

DNA replication occurs during S phase of the cell cycle (see figure 2.13). When DNA replicates, it unwinds, breaks, builds a new nucleotide chain, and mends (**figure 9.15**). Enzymes called helicases unwind and hold apart replicating DNA, enabling other enzymes to guide the assembly of a new DNA strand.

Human DNA replicates about 50 bases per second. To get the job done, a human chromosome replicates simultaneously at hundreds of points along its length, and the pieces join. A site where DNA is locally opened, resembling a fork, is called a **replication fork**.

DNA replication begins when a helicase breaks the hydrogen bonds that connect a base pair (**figure 9.16**). Binding proteins hold the two strands apart. Another enzyme, primase, then attracts complementary RNA nucleotides to build a short piece of RNA, called an RNA primer, at the start of each segment of DNA to be replicated. The RNA primer is required because the major replication enzyme, **DNA polymerase** (DNAP), can only add bases to an existing nucleic acid strand. (A polymerase is an enzyme that builds a polymer, which is a chain of chemical building blocks.) Next, the RNA primer attracts DNAP, which brings in DNA nucleotides complementary to the exposed bases on the parental strand; this strand serves as a mold, or template. New bases are added one at a time, starting at the RNA primer.

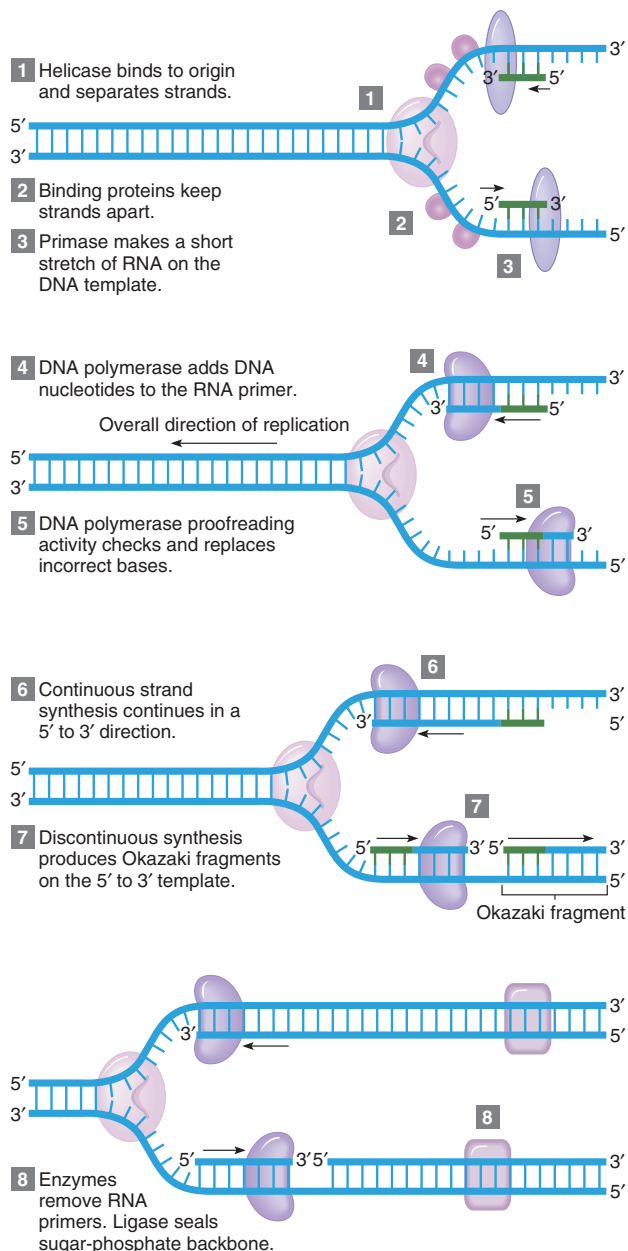
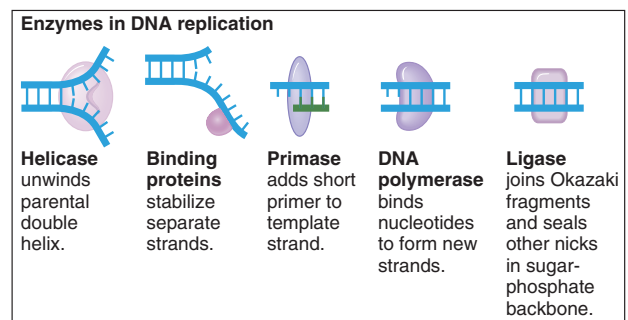


Figure 9.16 Activities at the replication fork. DNA replication takes many steps.

The new DNA strand grows as hydrogen bonds form between the complementary bases. The nucleotides are abundant in cells, and are synthesized from dietary nutrients.

DNAP works directionally, adding new nucleotides to the exposed 3' end of the sugar in the growing strand. Overall, replication proceeds in a 5' to 3' direction, because this is the only chemical configuration in which DNAP can add bases. How can the growing fork proceed in one direction, when both parental strands must be replicated? The answer is that on at least one strand, replication is discontinuous. It is accomplished in small pieces from the inner part of the fork outward, in a pattern similar to backstitching. Next, an enzyme called a **ligase** then seals the sugar-phosphate backbones of the pieces, building the new strand. These pieces, up to 150 nucleotides long, are called Okazaki fragments, after their discoverer (see figure 9.16).

DNA polymerase also “proofreads” as it goes, excising mismatched bases and inserting correct ones. It also removes the RNA primer and replaces it with the correct DNA bases. Yet another enzyme, called an annealing helicase, rewinds any sections of the DNA molecule that remain unwound. Finally, ligases seal the entire sugar-phosphate backbone. Ligase comes from a Latin word meaning “to tie.”

As a human body grows to 100 trillion or so cells, DNA replication occurs about 100 quadrillion times. Some of the

DNA sequence encodes protein, but most does not. The genome is in this sense a little like a booklet describing the parts of a machine, such as a car, that comes with a much more extensive manual that explains the details of keeping it in working order. The next chapters in this part of the book explain how the human instruction manual is accessed, and explores some of the nuances of its use. The *Bioethics: Choices for the Future* box discusses a controversial use of DNA testing.

Key Concepts

1. Experiments that followed the distribution of labeled DNA showed that DNA replication is semiconservative, not conservative or dispersive.
2. Enzymes replicate DNA.
3. DNA replication occurs simultaneously at several points on each chromosome, and the pieces join.
4. At each initiation site, primase directs synthesis of a short RNA primer, which DNA eventually replaces. DNA polymerase adds complementary bases to the RNA primer. Ligase joins the sugar-phosphate backbone.
5. DNA is synthesized in a 5' to 3' direction, discontinuously on one strand.

Bioethics: Choices for the Future

Infidelity Testing

“Afraid your significant other is cheating? Send us a DNA sample, and we’ll find the proof.”

Bridgette came home a day early from a business trip to find her husband Roy drinking coffee in their kitchen with Tiffany, his business associate. They were laughing so hard that it took a few moments for them to notice Bridgette standing there. When they did, Tiffany blushed and Roy knocked over her coffee mug, then they both stammered that they were discussing an acquisition. Bridgette didn’t buy it.

She went upstairs to unpack. Flinging her purse on the bed, she noticed several strands of brown hair on her pillow. Bridgette’s hair was brown, too, but she never left it on her pillow like that. She also noticed a crumpled tissue on the floor, part-way under the bed.

Bridgette knew just what to do. She’d recently read an article about companies that test “abandoned DNA,” so she went back downstairs for some plastic bags, and nonchalantly picked up Tiffany’s coffee mug, carrying it all back upstairs. In the bedroom, she quickly collected her evidence—the telltale hairs, the discarded used tissue, and on a cotton swab she rubbed along the inside rim of the mug. She e-mailed gotchaDNA.com and received a cheek swab collection kit a few days later, which she used to send in her own DNA for comparison, plus the \$600 fee. Then she waited.

The technicians at gotchaDNA.com extracted the DNA from the samples. First they checked for Y chromosome markers, found on the crumpled tissue. Then they looked for several STR (short tandem repeats—see table 5.3) markers and found what Bridgette

had feared—the DNA on the mug that Tiffany had used and in the hair cells matched each other, and not Bridgette’s DNA. Tiffany, or at least her hair, had somehow found its way onto Bridgette’s pillow.

Cells use DNA to manufacture protein. People use DNA to identify people. Chapter 1 introduced uses of DNA testing in several settings. Another is “infidelity DNA testing,” which dozens of companies offer on the Internet. Although a few websites provide documents for attesting that the samples are given willingly, many do not, and even list suggested sources of DNA for “adultery tracing.” These sources include underwear, toothbrushes, dental floss, nail clippings, gum, cigarette butts, and razor clippings.

Questions for Discussion

1. In the United Kingdom, a law was enacted to prohibit sampling of celebrity’s DNA after someone tried to steal hair from Prince Harry to determine whether or not Prince Charles is his biological father. The United States has no such law. Do you think that one is warranted? (A half waffle whose other half was consumed by Barack Obama was auctioned on eBay, with claims that it contained the presidential DNA.)
2. Do you think that DNA data obtained without consent should be admissible in a court of law? State a reason for your answer.
3. Discuss one reason in support of infidelity testing of DNA and one reason against it.
4. Identify the individuals in the scenario whom you believe behaved unethically.

Summary

9.1 Experiments Identify and Describe the Genetic Material

1. DNA encodes information that the cell uses to synthesize protein. DNA can also replicate, passing on its information.
2. Many experimenters described DNA as the hereditary material. Miescher identified DNA in white blood cell nuclei. Garrod connected heredity to enzyme abnormalities. Griffith identified a “transforming principle” that transmitted infectiousness in pneumonia-causing bacteria; Avery, MacLeod, and McCarty discovered that the transforming principle is DNA; and Hershey and Chase confirmed that the genetic material is DNA and not protein.
3. Levene described the three components of a DNA building block and found that they appear in DNA in equal amounts. Chargaff discovered that the amount of **adenine** (A) equals the amount of **thymine** (T), and the amount of **guanine** (G) equals that of **cytosine** (C). A and G are **purines**; C and T are **pyrimidines**. Rosalind Franklin showed that the molecule is a certain type of helix. Watson and Crick deduced DNA’s structure.

9.2 DNA Structure

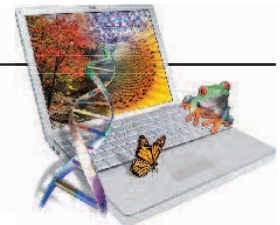
4. A **nucleotide** is a DNA building block. It consists of a **deoxyribose**, a phosphate, and a nitrogenous base.
5. The rungs of the DNA double helix consist of hydrogen-bonded **complementary base pairs** (A with T, and C with G). The rails are chains of alternating sugars and phosphates that run **antiparallel** to each other. DNA is highly coiled, and complexed with protein to form **chromatin**.

9.3 DNA Replication—Maintaining Genetic Information

6. Meselson and Stahl demonstrated the **semiconservative** nature of DNA replication with density shift experiments.
7. During replication, the DNA unwinds locally at several sites. **Replication forks** form as hydrogen bonds break between base pairs. Primase builds short RNA primers, which DNA sequences eventually replace. Next, **DNA polymerase** fills in DNA bases, and **ligase** seals the sugar-phosphate backbone.
8. Replication proceeds in a 5’ to 3’ direction, so the process must be discontinuous in short stretches on one strand.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

1. List the components of a nucleotide.
2. How does a purine differ from a pyrimidine?
3. DNA specifies and regulates the cell’s synthesis of protein. If a cell contains all the genetic material required to carry out protein synthesis, why must its DNA be replicated?
4. Why would a DNA structure in which each base type could form hydrogen bonds with any of the other three base types not produce a molecule that is easily replicated?
5. What part of the DNA molecule encodes information?
6. Explain how DNA is a directional molecule in a chemical sense.
7. Match the experiment described in the left column to a concept it illustrates in the right column (more than one answer may be possible).

<ol style="list-style-type: none"> 1. Density shift experiments 2. Discovery of an acidic substance that includes nitrogen and phosphorus on dirty bandages 3. “Blender experiments” that showed that the part of a virus that infects bacteria contains phosphorus, but not sulfur 4. Determination that DNA contains equal amounts of guanine and cytosine, and of adenine and thymine 5. Discovery that bacteria can transfer a “factor” that transforms a harmless strain into a lethal one 	<ol style="list-style-type: none"> a. DNA is the hereditary material b. Complementary base pairing is part of DNA structure and maintains a symmetrical double helix c. Identification of nuclein d. DNA, not protein, is the hereditary material e. DNA replication is semiconservative, not conservative or dispersive
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8. Place the following enzymes in the order in which they begin to function in DNA replication: ligase, primase, helicase and DNA polymerase
9. How can incredibly long DNA molecules fit into a cell’s nucleus?

- Place in increasing size order: nucleosome, histone protein, and chromatin.
- How are very long strands of DNA replicated without twisting into a huge tangle?
- List the steps in DNA replication.
- Why must DNA be replicated continuously as well as discontinuously?
- How does RNA participate in DNA replication?
- Describe two experiments that supported one hypothesis while also disproving another.
- Is downloading a document from the Internet analogous to replicating DNA? Cite a reason for your answer.

Applied Questions

- In Bloom syndrome, ligase malfunctions. As a result, replication forks move too slowly. Why?
- DNA contains the information that a cell uses to synthesize a particular protein. How do proteins assist in DNA replication?
- A person with deficient or abnormal ligase may have an increased cancer risk and chromosome breaks that cannot heal. The person is, nevertheless, alive. Why are there no people who lack DNA polymerase?
- Write the sequence of a strand of DNA replicated from each of the following base sequences:
 - TCGAGAATCTCGATT
 - CCGTATAGCCGGTAC
 - ATCGGATCGCTACTG
- Which do you think was the more far-reaching accomplishment, determining the structure of DNA, or sequencing the human genome? State a reason for your answer.
- Describe a recent news event, feature article, film, or television program that mentions a DNA sequence.
- Cite an example of how knowing a DNA sequence could be abused, and an example of how knowing a DNA sequence could be helpful.
- People often use the phrase “the gene for” to describe traits that do not necessarily or directly arise from a protein’s actions, such as “a gene for jealousy” or “a gene for acting.” How would you explain to them what a gene actually is?

Web Activities

- The Frozen Ark project is an international consortium of zoos, laboratories, and museums that is preserving DNA samples from endangered animal species. Consult <http://www.frozenark.org>
 - Follow one of the links and describe an endangered species. What do you think is the value of this project?
 - Do you think the project should be extended to include organisms other than animals? Cite a reason for your answer.
 - What would be the difficulties encountered in attempting to increase population sizes of endangered species using stored DNA?
- Visit the Cystic Fibrosis Mutation Database website. Select twenty contiguous bases of the sequence for the cystic fibrosis gene and write the complementary sequence.

Case Studies and Research Results

- Researchers at the University of Rochester studied children born with an infection called roseola. The children also have the causative herpes virus inserted into their chromosomes. At least one parent of each child had the virus in a chromosome of a sampled hair cell.
 - How did the children likely become infected?
 - What does the transmission of the viral DNA from generation to generation reveal about the structure of DNA and its replication?



The genetic code is universal. In a hippo and a human, the same mRNA triplets specify the same amino acids—that is, the genetic code rules are the same. The genome sequences of hippo and human, however, differ.

CHAPTER

10

Gene Action: From DNA to Protein

Chapter Contents

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RNA Structure and Types

Transcription Factors

Steps of Transcription

RNA Processing

10.3 Translation of a Protein

Deciphering the Genetic Code

Building a Protein

10.4 Processing a Protein

Protein Folding

Protein Misfolding

The Genetic Code Is Universal

A human might not appear to have much in common with a hippo, but we share the same genetic code—that is, the correspondence between the DNA sequence of a gene and the amino acid sequence of a protein. The “words” of the language of genes are three-base mRNA units called codons. A particular codon specifies the same amino acid, whether it is in a bacterium, a mushroom, a tree, or a mongoose. All life on Earth uses the same genetic code.

The universality of the genetic code has important consequences for medical technology. It means that we can use organisms other than ourselves to study diseases. We can put the gene that encodes human insulin into bacteria, which then read the gene and produce the protein product, insulin—used to treat people with diabetes. The universality of the genetic code also means that the genetic instructions for a protein that makes a jellyfish glow green can be placed next to any gene, in any organism, and the resulting fluorescent signal can be used to track the activity of that gene. Figure 19.1 shows mice displaying their jellyfish genes.

Many articles and reports, however, talk about the “human genetic code.” *“The effects of modern living may be damaging the human genetic code,”* reads one headline. The confusion arises from dual definitions of the word “code.” The primary definition is “a system of signals used to represent letters or numbers in transmitting messages,” such as a health code, building code, or moral code. However, “code” also refers to the sequence of symbols themselves, as in “computer code.”

When people say “my genetic code” or “the human genetic code,” what they generally mean is the genome sequence. This chapter explores how the genetic code was deciphered, and the remaining two chapters in this unit offer many examples of gene sequences.

10.1 From DNA to Protein

DNA replication preserves genetic information by giving new cells complete sets of operating instructions. Historically, molecular genetics has focused on the DNA sequences that encode protein—the genes. Yet, even before the genome was sequenced, researchers suspected that our DNA does much more than provide blueprints for proteins. Much of the human genome instead *controls* the protein-encoding genes, including when particular proteins are synthesized, at what rates, and in which cells under which conditions. These control sequences are abundant, lying within protein-encoding genes as well as among the vast expanses of DNA that do not encode protein. This chapter explains how DNA specifies proteins, and chapter 11 delves into some of the complexities of gene expression that sequencing the genome revealed.

Our genes encode, at last count, 20,325 proteins. A protein consists of one or more long chains of amino acids called polypeptides. A short sequence of amino acids is called a peptide, and the bonds that join amino acids are called peptide bonds. Proteins have a great variety of functions (**table 10.1**). They transport iron in the blood; provide immunity; digest food; clot blood; regulate use of glucose; forms skin, hair, and muscle; and enable cells to move.

To use the genetic information in the nucleus to synthesize proteins, the process of **transcription** first makes a copy of a gene that is an RNA molecule complementary to one strand of the DNA double helix. The RNA copy is taken out of the nucleus and into the cytoplasm. There, the process of **translation** uses the information in the RNA to manufacture a

protein by aligning and joining specified amino acids. Finally, the protein must fold into a specific three-dimensional form in order to function.

Accessing the genome is a huge, ongoing task. Cells replicate their DNA only during S phase of the cell cycle. In contrast, transcription and translation occur continuously, except during M phase. Transcription and translation supply the proteins essential for life, as well as those that give a cell its specialized characteristics.

10.2 Transcription

Watson and Crick, shortly after publishing their structure of DNA in 1953, described the relationship between nucleic acids and proteins as a directional flow of information called the “central dogma” (**figure 10.1**). As Francis Crick explained in 1957, “*The specificity of a piece of nucleic acid is expressed solely by the sequence of its bases, and this sequence is a code for the amino acid sequence of a particular protein.*” This statement inspired more than a decade of intense research to identify the participants in protein synthesis and discover how they interact. The process centers around RNA.

RNA is the bridge between gene and protein. RNA and DNA share an intimate relationship, as **figure 10.2** depicts. The bases of an RNA sequence are complementary to those of one strand of the double helix, which is called the **template strand**. An enzyme, **RNA polymerase**, assists the construction of an

Table 10.1 Protein Diversity in the Human Body

Protein	Function
Actin, myosin, dystrophin	Muscle contraction
Antibodies, antigens, cytokines	Immunity
Carbohydrases, lipases, proteases, nucleases	Digestion (digestive enzymes)
Casein	Milk protein
Collagen, elastin, fibrillin	Connective tissue
Colony-stimulating factors, erythropoietin	Blood cell formation
DNA and RNA polymerase	DNA replication, gene expression
Ferritin	Iron transport in blood
Fibrin, thrombin	Blood clotting
Growth factors, kinases, cyclins	Cell division
Hemoglobin, myoglobin	Oxygen transport
Insulin, glucagon	Control of blood glucose level
Keratin	Hair structure
Tubulin, actin	Cell movements
Tumor suppressors	Cancer prevention

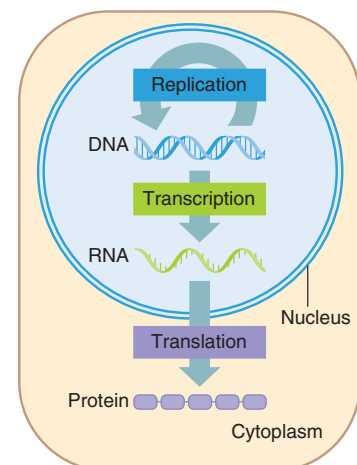


Figure 10.1 DNA to RNA to protein. Some of the information stored in DNA is copied to RNA (transcription), some of which is used to assemble amino acids into proteins (translation). DNA replication perpetuates genetic information. This figure repeats throughout the chapter, with the part under discussion highlighted.

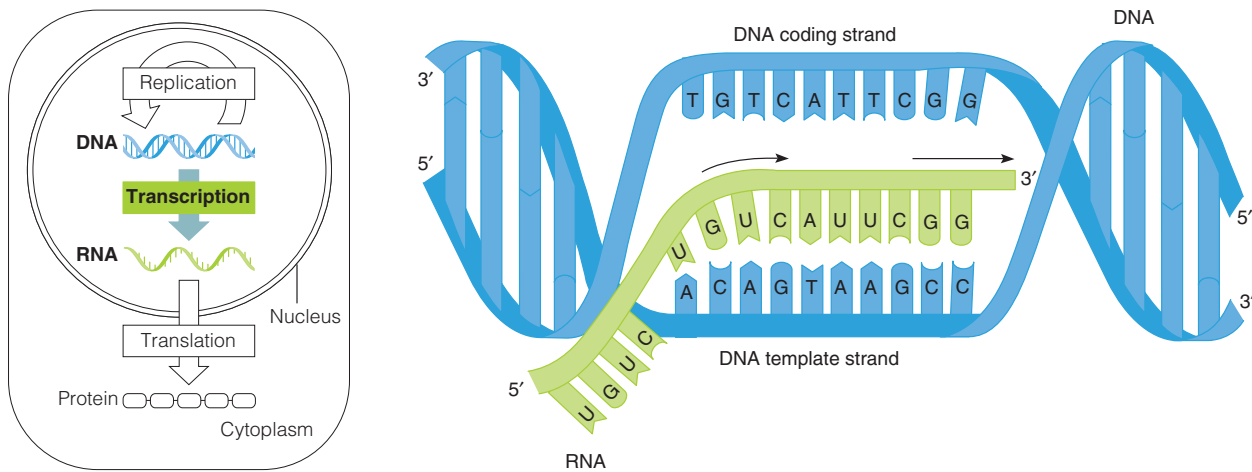


Figure 10.2 The relationship among RNA, the DNA template strand, and the DNA coding strand. The RNA sequence is complementary to the DNA template strand. This is the same sequence as the DNA coding strand, with uracil (U) in place of thymine (T).

RNA molecule. The other strand of the DNA double helix is called the **coding strand**.

DNA

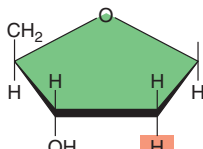
Stores RNA- and protein-encoding information, and transfers information to daughter cells

a.



Double-stranded

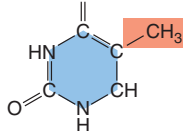
b.



Deoxyribose as the sugar

c.

Bases used:



Thymine (T)
Cytosine (C)
Adenine (A)
Guanine (G)

d.

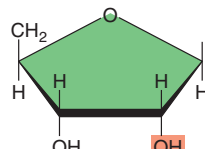
RNA

Carries protein-encoding information, and helps to make proteins

a.



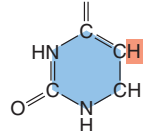
Generally single-stranded



Ribose as the sugar

b.

Bases used:



Uracil (U)
Cytosine (C)
Adenine (A)
Guanine (G)

c.

RNA Structure and Types

RNA and DNA have similarities and differences (**figure 10.3** and **table 10.2**). Both are nucleic acids, consisting of sequences of nitrogen-containing bases joined by sugar-phosphate backbones. However, RNA is usually single-stranded, whereas DNA is double-stranded. Also, RNA has the pyrimidine base **uracil** where DNA has thymine. As their names imply, RNA nucleotides include the sugar ribose, rather than DNA's deoxyribose. Functionally, DNA stores genetic information, whereas RNA controls how that information is used. The presence of the —OH at the 5' position of ribose makes RNA much less stable than DNA, which is critical in its function as a short-lived carrier of genetic information.

As RNA is synthesized along DNA, it folds into a three-dimensional shape, or **conformation**, that is determined by complementary base pairing within the same RNA molecule. For example, a sequence of AAUUUCC might hydrogen bond

Table 10.2 How DNA and RNA Differ

DNA	RNA
1. Usually double-stranded	1. Usually single-stranded
2. Thymine as a base	2. Uracil as a base
3. Deoxyribose as the sugar	3. Ribose as the sugar
4. Maintains protein-encoding information	4. Carries protein-encoding information and controls how information is used
5. Cannot function as an enzyme	5. Can function as an enzyme
6. Persists	6. Transient

Figure 10.3 DNA and RNA differences. (a) DNA is double-stranded; RNA is usually single-stranded (b). DNA nucleotides include deoxyribose, whereas RNA nucleotides have ribose (c). Finally, DNA nucleotides include the pyrimidine thymine, whereas RNA has uracil (d).

Type of RNA	Size (number of nucleotides)	Function
mRNA	500–4,500 +	Encodes amino acid sequence
rRNA	100–3,000	Associates with proteins to form ribosomes, which structurally support and catalyze protein synthesis
tRNA	75–80	Transports specific amino acids to the ribosome for protein synthesis

to a sequence of UAAAAGG—its complement—elsewhere in the same molecule, a little like touching elbows to knees. These shapes are very important for RNA’s functioning. The three major types of RNA are messenger RNA, ribosomal RNA, and transfer RNA (table 10.3). Many of these other classes of RNA control which genes are expressed (transcribed and translated) under specific circumstances. Table 11.4 describes other types of RNA molecules.

Messenger RNA (mRNA) carries the information that specifies a particular protein. Each three mRNA bases in a row form a genetic code word, or **codon**, that specifies a certain amino acid. Because genes vary in length, so do mature mRNA molecules. Most mRNAs are 500 to 4,500 bases long. Differentiated cells can carry out specialized functions because they express certain subsets of genes—that is, they produce certain mRNA molecules, also called transcripts. The information in the transcripts is then used to manufacture the encoded proteins. A muscle cell, for example, has many mRNAs that specify the contractile proteins actin and myosin, whereas a skin cell contains many mRNAs that specify the scaly keratin proteins.

To use the information in an mRNA sequence, a cell requires the two other major classes of RNA. **Ribosomal RNA (rRNA)** molecules range from 100 to nearly 3,000 nucleotides long. Ribosomal RNAs associate with certain proteins to form a ribosome. Recall from chapter 2 that a ribosome is an organelle made up of many different protein and RNA subunits. Overall, a ribosome functions as a machine to attach amino acids to form proteins (figure 10.4).

A ribosome has two subunits that are separate in the cytoplasm but join at the site of initiation of protein synthesis. The larger ribosomal subunit has three types of rRNA molecules, and the small subunit has one. Ribosomal RNA, however, is

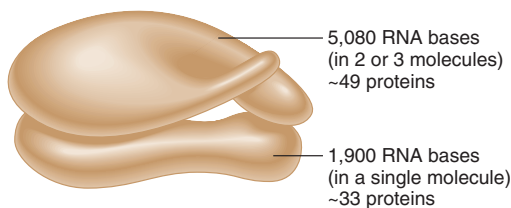


Figure 10.4 The ribosome. A ribosome from a eukaryotic cell has two subunits; together, they consist of 82 proteins and four rRNA molecules.

more than a structural support. Certain rRNAs catalyze the formation of the peptide bonds between amino acids. Such an RNA with enzymatic function is called a ribozyme. Other rRNAs help to align the ribosome and mRNA.

The third major type of RNA molecule, **transfer RNA (tRNA)**, binds an mRNA codon at one end and a specific amino acid at the other. A tRNA molecule is only 75 to 80 nucleotides long. Some of its bases form weak chemical bonds with each other, folding the tRNA into loops in

a characteristic cloverleaf shape (figure 10.5). One loop of the tRNA has three bases in a row that form the **anticodon**, which is complementary to an mRNA codon. The end of the tRNA opposite the anticodon strongly bonds to a specific amino acid. A tRNA with a particular anticodon sequence always carries the same amino acid. (Organisms have 20 types of amino acids.) For example, a tRNA with the anticodon sequence GAA always picks up the amino acid phenylalanine. Enzymes attach amino acids to tRNAs that bear the appropriate anticodons, where they form chemical bonds (figure 10.6).

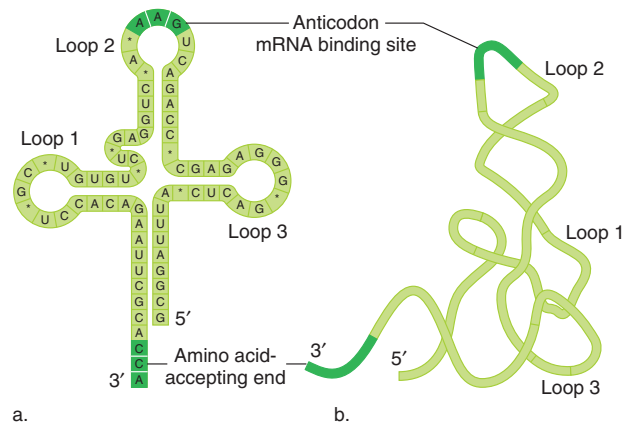


Figure 10.5 Transfer RNA. (a) Certain nucleotide bases within a tRNA hydrogen bond with each other to give the molecule a “cloverleaf” conformation that can be represented in two dimensions. The darker bases at the top form the anticodon, the sequence that binds a complementary mRNA codon. Each tRNA terminates with the sequence CCA, where a particular amino acid covalently bonds. A three-dimensional representation of a tRNA (b) depicts the loops that interact with the ribosome.

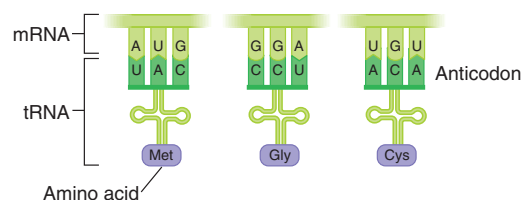


Figure 10.6 A tRNA with a particular anticodon sequence always binds the same type of amino acid.

Transcription Factors

Study of the control of gene expression began in 1961, when French biologists François Jacob and Jacques Monod described the remarkable ability of *E. coli* bacteria to produce the enzymes to metabolize the sugar lactose—but only when lactose is in the cell's surroundings. What “tells” a simple bacterial cell to transcribe the proteins it needs, at exactly the right time?

Jacob and Monod discovered that a modified form of lactose “turned on” the genes whose encoded proteins break it down. They named the set of genes that are coordinately controlled an operon, writing in 1961, “*The genome contains not only a series of blueprints, but a coordinated program of protein synthesis and means of controlling its execution.*”

In bacteria, operons turn transcription of a few genes on or off. In more complex organisms, different cell types express different subsets of genes. To manage this, groups of proteins called **transcription factors** come together, forming an apparatus that binds DNA at certain sequences and initiates transcription at specific sites on chromosomes. (Bacterial regulatory proteins are also called transcription factors.) The transcription factors, activated by signals from outside the cell such as hormones and growth factors, set the stage for transcription by forming a pocket for RNA polymerase—the enzyme that builds an RNA chain.

Transcription factors include regions called binding domains that guide them to the genes they control. The DNA binding domains have very colorful names, such as “helix-turn-helix,” “zinc fingers,” and “leucine zippers,” that reflect their distinctive shapes.

The human genome encodes about 2,600 proteins that have DNA binding domains and about 2,000 of these are thought to function as transcription factors. There are only a few classes of transcription factors, however, based on similar overall structures. The fact that transcription factors work in combinations provides great specificity in controlling gene expression. Overall, transcription factors link the genome to the environment. For example, lack of oxygen, such as from choking or smoking, sends signals that activate transcription factors to turn on dozens of genes that enable cells to handle the stress of low-oxygen conditions.

Mutations in transcription factor genes can have wide-ranging effects, because the factors control many genes. The varied symptoms of Rett syndrome, discussed in Reading 6.2, arise from a mutation in the gene that encodes a transcription factor called MECP2. Transcription factors are themselves controlled by each other and by other classes of molecules.

Steps of Transcription

Transcription and translation are each described in three steps: initiation, elongation, and termination.

How do transcription factors and RNA polymerase (RNAP) “know” where to bind to DNA to begin transcribing a specific gene? In transcription initiation, transcription factors and RNA polymerase are attracted to a **promoter**, which is a special sequence that signals the start of the gene. Signals

from outside the cell alter the chromatin structure in a way that exposes the promoter of a gene whose transcription is required under the particular conditions (see figure 11.8).

Figure 10.7 shows a simplified view of transcription factor binding, which sets up a site called a preinitiation complex to receive RNA polymerase. The first transcription factor to bind, called a TATA binding protein, is chemically attracted to a DNA sequence called a TATA box—the base sequence TATA surrounded by long stretches of G and C. Once the first transcription factor binds, it attracts others in groups. Finally RNA polymerase joins the complex, binding just in front of the start of the gene sequence. The assembly of these components is transcription initiation.

In the next stage, transcription elongation, enzymes unwind the DNA double helix locally, and free RNA nucleotides bond with exposed complementary bases on the DNA template strand (see figure 10.2). RNA polymerase adds the RNA nucleotides in the sequence the DNA specifies, moving along the DNA strand in a 3' to 5' direction, synthesizing the RNA molecule in a 5' to 3' direction. A terminator sequence in the DNA indicates where the gene's RNA-encoding region ends. When this spot is reached, the third stage, transcription termination, occurs (**figure 10.8**). A typical rate of transcription in humans is 20 bases per second.

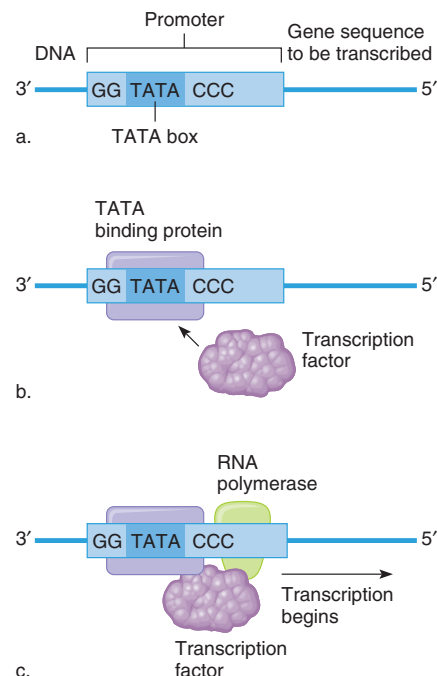


Figure 10.7 Setting the stage for transcription to begin. (a) Proteins that initiate transcription recognize specific sequences in the promoter region of a gene. (b) A binding protein recognizes the TATA region and binds to the DNA. This allows other transcription factors to bind. (c) The bound transcription factors form a pocket that allows RNA polymerase to bind and begin making RNA.

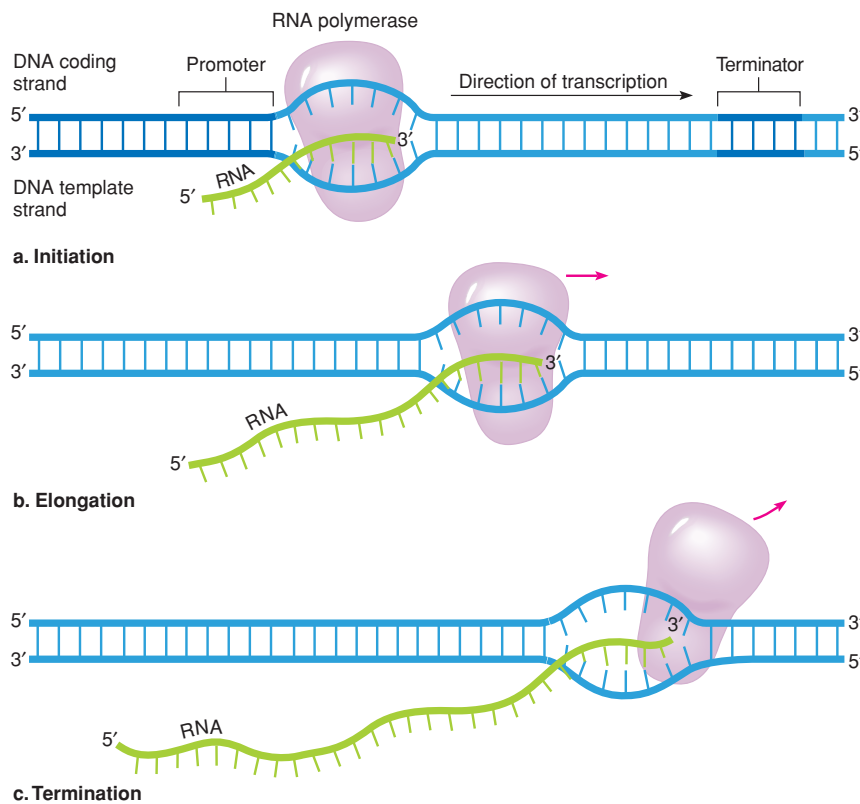


Figure 10.8 Transcription of RNA from DNA. Transcription occurs in three stages: initiation, elongation, and termination. Initiation is the control point that determines which genes are transcribed. RNA nucleotides are added during elongation. A terminator sequence in the gene signals the end of transcription.

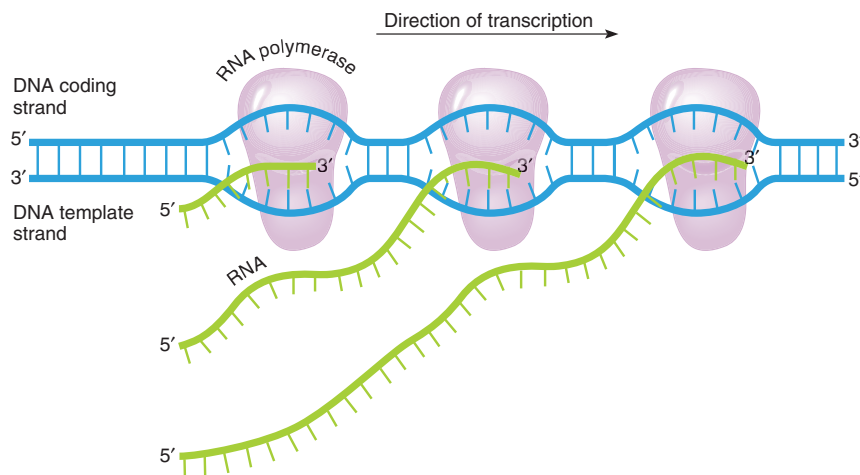


Figure 10.9 Many identical copies of RNA are transcribed simultaneously. Usually 100 or more DNA bases lie between RNA polymerases.

RNA is typically transcribed using only a gene's template strand. However, different genes on the same chromosome may be transcribed from different strands of the double helix. The coding strand of the DNA is so-called because its sequence is identical to that of the RNA, except with thymine (T) in place of uracil (U). Several RNAs may be transcribed from the same DNA template strand simultaneously due to the —OH at the 2' position in ribose. (figure 10.9) Since mRNA is short-lived, with about half of it degraded every 10 minutes, a cell must constantly transcribe certain genes to maintain supplies of essential proteins.

To determine the sequence of RNA bases transcribed from a gene, write the RNA bases that are complementary to the template DNA strand, using uracil opposite adenine. For example, a DNA template strand that has the sequence

C C T A G C T A C

is transcribed into RNA with the sequence

G G A U C G A U G

and the coding DNA sequence is

G G A T C G A T G.

RNA Processing

In bacteria, RNA is translated into protein as soon as it is transcribed from DNA because a nucleus does not physically separate the two processes. In eukaryotic cells, mRNA must first exit the nucleus to enter the cytoplasm, where ribosomes are located. Messenger RNA is altered before it is translated in these more complex cells. The modifications take several steps.

First, after mRNA is transcribed, a short sequence of modified nucleotides, called a cap, is added to the 5' end of the molecule. The cap consists of a backwardly inserted guanine (G), which attracts an enzyme that adds methyl groups (CH₃) to the G and one or two adjacent nucleotides. This methylated cap is a recognition site for protein synthesis. At the 3' end, a special polymerase adds about 200 adenines, forming a "poly A tail." The poly A tail is necessary for protein synthesis to begin, and may also stabilize the mRNA so that it stays intact longer.

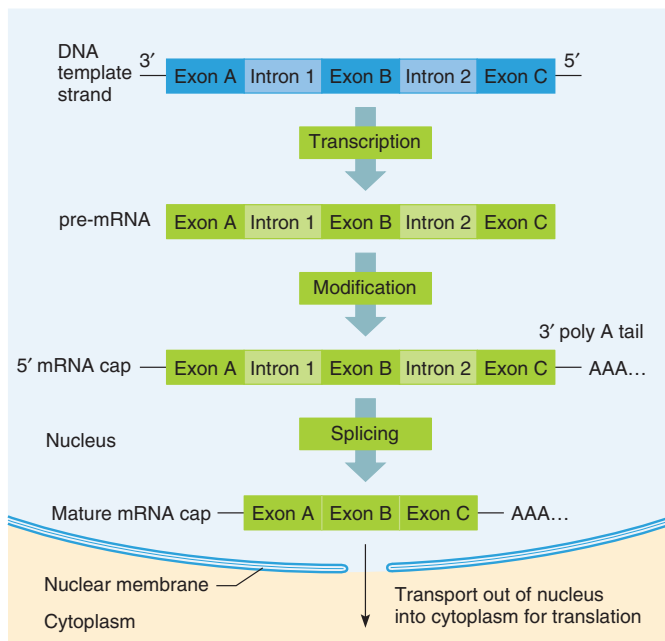


Figure 10.10 Messenger RNA processing—the maturing of the message. Several steps process pre-mRNA into mature mRNA. First, a large region of DNA containing the gene is transcribed. Then a modified nucleotide cap and poly A tail are added and introns are spliced out. Finally, the intact, mature mRNA is sent out of the nucleus.

Further changes occur to the capped, poly A tailed mRNA before it is translated into protein. Parts of mRNAs called **introns** (short for “intervening sequences”) that were transcribed are removed. The ends of the remaining molecule are spliced together before the mRNA is translated. The parts of mRNA that remain and are translated are called **exons** (figure 10.10).

Once introns are spliced out, enzymes check, or proofread, the remaining mRNA. Messenger RNAs that are too short or too long may be held in the nucleus. Proofreading also monitors tRNAs, ensuring that they assume the correct conformation.

Prior to intron removal, the mRNA is called pre-mRNA. Introns control their own removal. They associate with certain proteins to form small nuclear ribonucleoproteins (snRNPs), or “snurps.” Four snurps form a structure called a spliceosome that cuts introns out and attaches exons to form the mature mRNA that exits the nucleus. The introns cut themselves out of the RNA.

Introns range in size from 65 to 10,000 or more bases; the average intron is 3,365 bases. The average exon, in contrast, is only 145 bases long. The number, size, and organization of introns vary from gene to gene. The coding portion of the average human gene is 1,340 bases, whereas the average total size of a gene is 27,000 bases. The dystrophin gene is 2,500,000 bases, but its corresponding mRNA sequence is only 14,000 bases! The gene contains 80 introns.

The discovery of introns in the 1970s surprised geneticists, who had thought genes were like sentences in which all of the information has meaning. At first, some geneticists called introns “junk DNA”—a term that has unfortunately persisted even as researchers have discovered the functions of many introns. A few introns are known to encode some of the RNAs that control gene expression. Some introns are actually exons on the complementary strand of DNA. Introns may also be vestiges of ancient genes that have lost their original function, or are remnants of the DNA of viruses that once integrated into a chromosome.

The intron/exon organization of most genes provides a mechanism to maximize genetic information. Different combinations of exons of a particular gene encode different versions of the protein product, termed isoforms. From 40 to 60 percent of human genes encode isoforms, and the mechanism of combining exons in different ways is called **alternate splicing**.

Alternate splicing may explain how cell types use the same protein in slightly different ways in different tissues (see figure 11.9). For example, a protein that transports fats is shorter in the small intestine, where it carries dietary fats, than it is in the liver, where it carries fats made in the body.

Key Concepts

1. RNA is single-stranded, has uracil and ribose, and has different functions than DNA.
2. Messenger RNA transmits information to build proteins. Each three mRNA bases in a row form a codon that specifies a particular amino acid.
3. Ribosomal RNA and proteins form ribosomes, which physically support protein synthesis and help catalyze bonding between amino acids.
4. Transfer RNAs connect mRNA codons to amino acids.
5. Bacterial operons are simple gene control systems. In more complex organisms, cascades of transcription factors control gene expression.
6. RNA polymerase inserts complementary RNA bases opposite the DNA template strand.
7. Messenger RNA (mRNA) gains a modified nucleotide cap and a poly A tail.
8. Introns are transcribed and cut out, and exons are reattached. Introns are common and large in human genes.
9. Certain genes are processed into different-sized RNAs in different cell types.

10.3 Translation of a Protein

Transcription copies the information in DNA into the complementary language of RNA. The next step is translating mRNA into the precise sequence of amino acids that forms a protein. Particular mRNA codons correspond to particular amino acids

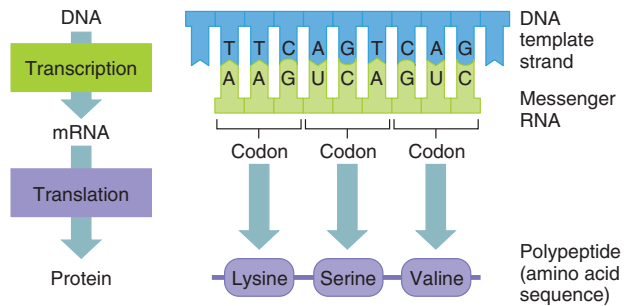


Figure 10.11 From DNA to RNA to protein. Messenger RNA is transcribed from a locally unwound portion of DNA. In translation, transfer RNA matches mRNA codons with amino acids.

(figure 10.11). This correspondence between the chemical languages of mRNA and protein is the **genetic code**. Translation occurs on free ribosomes in the cytoplasm as well as on ribosomes that are embedded in the endoplasmic reticulum (ER).

Francis Crick hypothesized that an “adaptor” molecule would enable the RNA message to attract and link amino acids into proteins. He envisioned “20 different kinds of adaptor molecule, one for each amino acid, and 20 different kinds of adaptor enzymes to join the amino acids to their adaptors.” In the 1960s, researchers deciphered the genetic code, determining which mRNA codons correspond to which amino acids. **Figure 10.12** displays the genetic code.

Deciphering the Genetic Code

The researchers who deciphered the genetic code used logic and experiments. More recently, annotation of the human genome sequence has confirmed and extended the earlier work, revealing new nuances in the genetic code. To understand how the genetic code works, it is helpful to ask the questions researchers asked in the 1960s.

Question 1—How Many RNA Bases Specify One Amino Acid?

The number of different protein building blocks (20) exceeds the number of different mRNA building blocks (4). Therefore, each codon must include more than one mRNA base. If a codon consisted of only one mRNA base, then codons could specify only four different amino acids, one corresponding to each of the four bases: A, C, G, and U. If each codon consisted of two bases, then only 16 (4^2) different amino acids could be specified, one corresponding to each of the 16 possible combinations of two RNA bases. If a codon consisted of three bases, then the genetic code could specify as many as 64 (4^3) different amino acids, sufficient to encode the 20 different amino acids that make up proteins. Therefore, the minimum number of bases in a codon is three.

Francis Crick and his coworkers experimented on a virus called T4 to confirm the triplet nature of the genetic code. They exposed the virus to chemicals that add or remove one, two, or three bases, and examined a viral gene with a well known

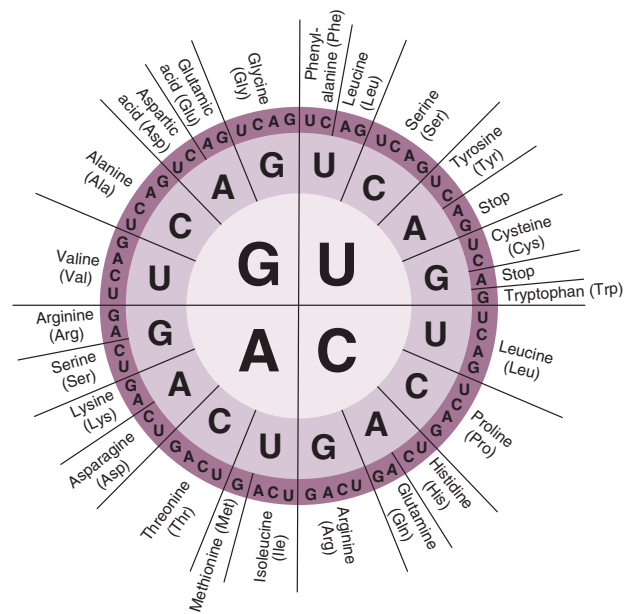


Figure 10.12 The genetic code. The first codon position is at the center of the circle. For example, the mRNA codon CUA encodes the amino acid leucine.

sequence and protein product. Altering the DNA sequence by one or two bases produced a different amino acid sequence. This happened because the change disrupted the **reading frame**, which is the sequence of amino acids encoded from a certain starting point in a DNA sequence. However, adding or deleting three contiguous bases added or deleted only one amino acid in the protein without disrupting the reading frame. The rest of the amino acid sequence was retained. The code, the researchers deduced, is triplet (**figure 10.13**).

Further experiments confirmed the triplet nature of the genetic code. Adding a base at one point in the gene and deleting a base at another point disrupted the reading frame only between these sites. The result was a protein with a stretch of the wrong amino acids, like a sentence with a few words in the middle that are misspelled.

Question 2—Does the Information in a DNA Sequence Overlap?

Consider a hypothetical mRNA sequence:

AUGCCCAAG

If the genetic code is triplet and a DNA sequence is “read” in a nonoverlapping manner, then this sequence has only three codons and specifies three amino acids:

AUGCCCAAG

AUG (methionine)

CCC (proline)

AAG (lysine)

Size of a genetic code word (codon)

Original RNA sequence	GAC GAC GAC GAC GAC GAC GAC ...
Amino acid sequence	Asp Asp Asp Asp Asp Asp Asp
One base added	GAC GGA CGA CGA CGA CGA CGA ...
Amino acid sequence altered	Asp Gly Arg Arg Arg Arg Arg
Two bases added	GAC UGG ACG ACG ACG ACG ...
Amino acid sequence altered	Asp Trp Thr Thr Thr Thr Thr
Three bases added	GAC UUG GAC GAC GAC GAC GAC ...
Amino acid sequence altered and then restored	Asp Leu Asp Asp Asp Asp Asp

 = Wrong triplet

Figure 10.13 Three at a time. Adding or deleting one or two nucleotides in a DNA sequence results in a frameshift that disrupts the encoded amino acid sequence. Adding or deleting three bases does not disrupt the reading frame because the code is triplet. This is a simplified representation of the Crick experiment.

If the DNA sequence is overlapping, however, the sequence specifies seven codons:

- AUGCCAAG
- AUG (methionine)
- UGC (cysteine)
- GCC (alanine)
- CCC (proline)
- CCA (proline)
- CAA (glutamine)
- AAG (lysine)

An overlapping DNA sequence seems to pack maximal information into a limited number of bases. However, certain amino acids would always follow certain others, constraining protein structure. For example, AUG would always be followed by an amino acid whose codon begins with UG. This does not happen. Therefore, the protein-encoding DNA sequence is not overlapping.

Even though the genetic code is non-overlapping, it is possible to read any DNA or RNA sequence in three different reading frames, depending upon the “start” base. **Figure 10.14** depicts the three possible reading frames for the sequence just

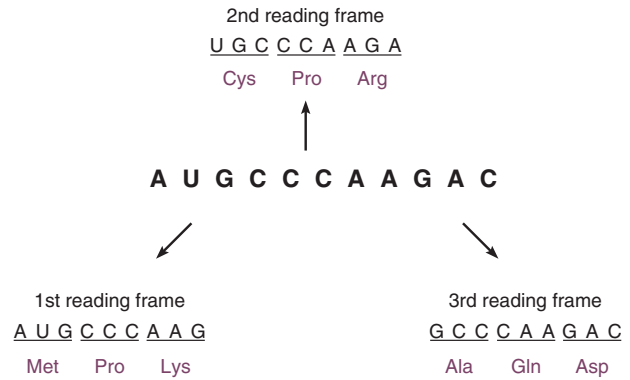


Figure 10.14 Reading frames—where the sequence begins. A sequence of DNA has three reading frames.

discussed, slightly extended. It encodes three different trios of amino acids.

Question 3—Can mRNA Codons Specify Anything Other Than Amino Acids?

Chemical analysis eventually showed that the genetic code includes directions for starting and stopping translation. The codon AUG signals “start,” and the codons UGA, UAA, and UAG signify “stop.” If the genetic code is compared to a sentence, the “start” codon is like the capital letter that begins the sentence, and the “stop” codon is like the period that ends the sentence. Another form of “punctuation,” a short sequence of bases at the start of each mRNA, enables the mRNA to form hydrogen bonds with rRNA in a ribosome. It is called a leader sequence.

Question 4—Do All Species Use the Same Genetic Code?

All species use the same mRNA codons to specify the same amino acids, as discussed in the chapter opener. The simplest explanation for the “universality” of the genetic code is that all life evolved from a common ancestor. No other mechanism as efficient at directing cellular activities has emerged and persisted.

The only known exceptions to the universality of the genetic code are a few codons in mitochondria and in certain single-celled eukaryotes (ciliated protozoa). These deviations may be tolerated because they do not affect the major repositories of DNA. The mitochondrial genome is small, and the affected ciliated protozoa have a second, smaller nucleus that houses some genes with one or two alternate codon-amino acid associations. In both cases, the major DNA sites adhere to the universal genetic code. Some types of single-celled organisms translate a stop codon into a twenty-first type of amino acid. Overall, however, the genetic code is considered to be universal.

Question 5—Which Codons Specify Which Amino Acids?

The number of bases in a codon, the non-overlapping reading frame, and universality are general features of the genetic code. The big question, back in the 1960s, was the code itself: Which codons specify which amino acids? In 1961, Marshall Nirenberg and Heinrich Matthaei at the National Institutes of Health used a precise and logical series of experiments to “crack the code.” They synthesized very simple mRNA molecules and added them to test tubes that contained all the chemicals and structures needed for translation, extracted from *E. coli* cells. Which amino acid would each synthetic RNA specify?

The first synthetic mRNA they made had the sequence UUUUUU. . . . In the test tube, this was translated into a peptide consisting entirely of one amino acid type: phenylalanine. This was the first entry in the genetic code dictionary: The codon UUU specifies the amino acid phenylalanine. The next experiments revealed that AAA codes for the amino acid lysine and CCC for proline. (GGG was unstable, so this part of the experiment could not be done.)

To reveal other codon-amino acid pairs, researchers synthesized chains of alternating bases. Synthetic mRNA of sequence AUAUUAU . . . introduced codons AUA and UAU. When translated, the mRNA yielded an amino acid sequence of alternating isoleucines and tyrosines. But was AUA the code for isoleucine and UAU for tyrosine, or vice versa? Another experiment with a more complex sequence answered the question.

The mRNA UUUUAUAUUUAUA, when translated from the first U of a UUU, encoded alternating phenylalanine and isoleucine. Because the first experiment had showed that UUU codes for phenylalanine, AUA must code for isoleucine. If AUA codes for isoleucine, then UAU must code for tyrosine (table 10.4).

By the end of the 1960s, researchers had used such logic to decipher the entire genetic code. Sixty of the possible 64

codons specify particular amino acids, three indicate “stop,” and one encodes both the amino acid methionine and “start.” This means that some amino acids are specified by more than one codon. For example, both UUU and UUC encode phenylalanine.

Different codons that specify the same amino acid are termed **synonymous codons**, just as synonyms are words with the same meaning. The genetic code is said to be degenerate because most amino acids are not uniquely specified. Synonymous codons often differ from one another by the base in the third position. The corresponding base of a tRNA’s anticodon is called the “wobble” position because it can bind to more than one type of base in synonymous codons. The degeneracy of the genetic code protects against mutation, because changes in the DNA that substitute a synonymous codon do not alter the protein’s amino acid sequence. **Nonsynonymous codons** encode different amino acids.

In the 1950s and 1960s, molecular genetics was still a very young science, and so the code breakers came mostly from the ranks of chemistry, physics, and math. Some of the more exuberant personalities organized an “RNA tie club” and inducted a member whenever someone added a piece to the puzzle of the genetic code, anointing him (there was no prominent “her”) with a tie and tie pin emblazoned with the structure of the specified amino acid.

The human genome project picked up where the genetic code experiments of the 1960s left off by identifying the DNA sequences that are transcribed into tRNAs. That is, 61 different tRNAs could theoretically exist, one for each codon that specifies an amino acid (the 64 triplets minus 3 stop codons). However, only 49 different genes encode tRNAs. This is because the same type of tRNA can detect synonymous codons that differ only in whether the wobble (third) position is U or C. The same type of tRNA, for example, binds to both UUU and UUC codons, which specify the amino acid phenylalanine. Synonymous codons ending in A or G use different tRNAs. Sequencing of other genomes reveals that some types of organisms preferentially use particular codons for amino acids specified by more than one type of codon. Researchers do not yet understand the significance, if any, of such “codon usage bias.”

Building a Protein

Protein synthesis requires mRNA, tRNA molecules carrying amino acids, ribosomes, energy-storing molecules such as adenosine triphosphate (ATP) and guanosine triphosphate (GTP), and various protein factors. These pieces meet in a stage called translation initiation (figure 10.15). Chemical bonds hold the different components together.

First, the mRNA leader sequence forms hydrogen bonds with a short sequence of rRNA in a small ribosomal

Table 10.4 Deciphering RNA Codons and the Amino Acids They Specify

Synthetic RNA	Encoded Amino Acid Chain	Puzzle Piece
UUUUUUUUUUUUUUUUUU	Phe-Phe-Phe-Phe-Phe-Phe	UUU = Phe
AAAAAAAAAAAAAAAAAA	Lys-Lys-Lys-Lys-Lys-Lys	AAA = Lys
GGGGGGGGGGGGGGGGGG	Gly-Gly-Gly-Gly-Gly-Gly	GGG = Gly
CCCCCCCCCCCCCCCCCC	Pro-Pro-Pro-Pro-Pro-Pro	CCC = Pro
AUAUUAUAUUAUUAUUAU	Ile-Tyr-Ile-Tyr-Ile-Tyr	AUA = Ile or Tyr UAU = Ile or Tyr
UUUAUAUUUAUUAUUUAU	Phe-Ile-Phe-Ile-Phe-Ile	AUA = Ile UAU = Tyr

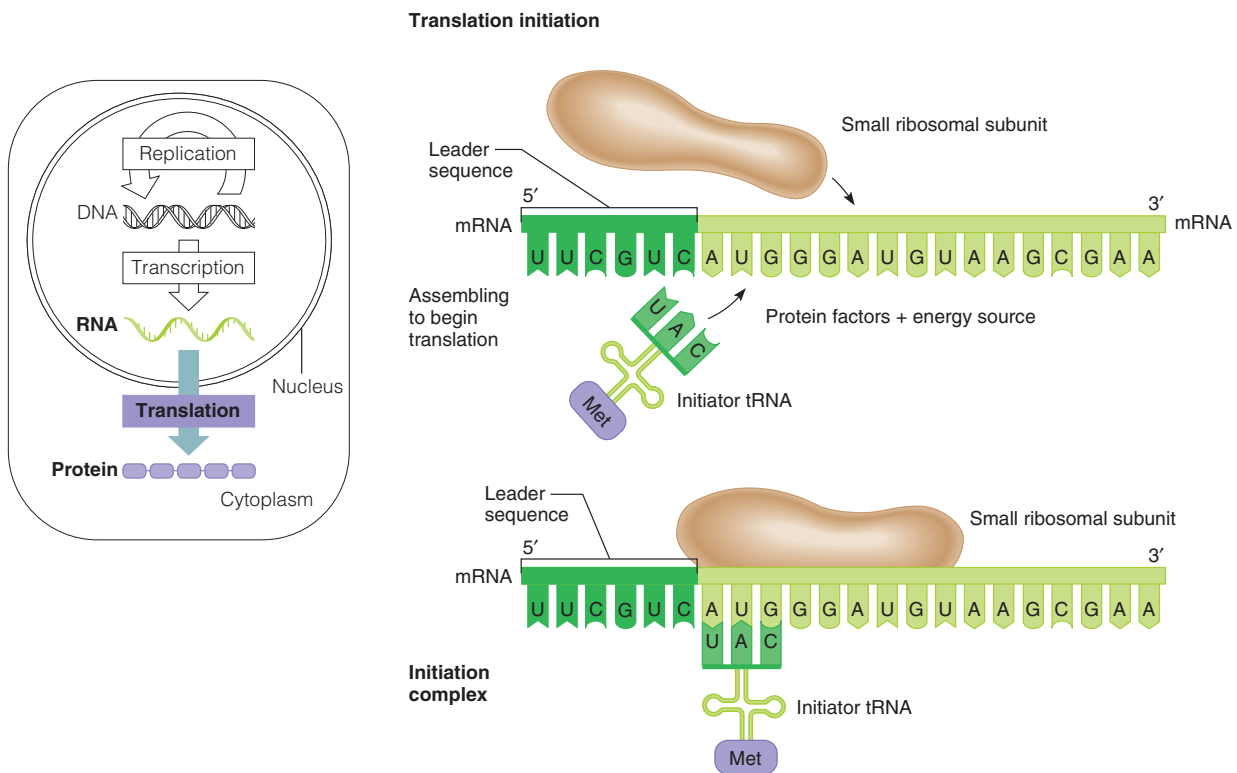


Figure 10.15 Translation begins as the initiation complex forms. Initiation of translation brings together a small ribosomal subunit, mRNA, and an initiator tRNA, and aligns them in the proper orientation to begin translation.

subunit. The first mRNA codon to specify an amino acid is always AUG, which attracts an initiator tRNA that carries the amino acid methionine (abbreviated *met*). This methionine signifies the start of a polypeptide. The small ribosomal subunit, the mRNA bonded to it, and the initiator tRNA with its attached methionine form the initiation complex at the appropriate AUG codon of the mRNA.

To start the next stage, elongation, a large ribosomal subunit bonds to the initiation complex. The codon adjacent to the initiation codon (AUG), which is GGA in **figure 10.16**, then bonds to its complementary anticodon, which is part of a free tRNA that carries the amino acid glycine. The two amino acids (*met* and *gly* in the example), still attached to their tRNAs, align.

The part of the ribosome that holds the mRNA and tRNAs together can be described as having two sites. The positions of the sites on the ribosome remain the same with respect to each other as translation proceeds, but they cover different parts of the mRNA as the ribosome moves. The P site holds the growing amino acid chain, and the A site right next to it holds the next amino acid to be added to the chain. In **figure 10.14**, when the forming protein consists of only the first two amino acids, *met* occupies the P site and *gly* the A site.

With the help of rRNA that functions as a ribozyme, the amino acids link by forming a specific type of chemical bond called a peptide bond. Then the first tRNA is released. It will pick up another amino acid of the same type and be used again. Special enzymes ensure that tRNAs always pick up the correct amino acids, which is crucial to the accuracy of translation. The ribosome and its attached mRNA are now bound to a single tRNA, with two amino acids extending from it at the P site. This is the start of a polypeptide.

Next, the ribosome moves down the mRNA by one codon. The region of the mRNA that was at the A site is thus now at the P site. A third tRNA enters at the now-vacated new A site, corresponding to the next codon, carrying its amino acid (*cys* in **figure 10.16b**). This third amino acid aligns with the other two and forms a peptide bond to the second amino acid in the growing chain, now extending from the P site. The tRNA attached to the second amino acid is released and recycled. The polypeptide continues to build, one amino acid at a time. Each piece is brought in by a tRNA whose anticodon corresponds to a consecutive mRNA codon as the ribosome moves down the mRNA (**figure 10.16c**).

Elongation halts when the A site of the ribosome has a “stop” codon (UGA, UAG, or UAA), because no tRNA

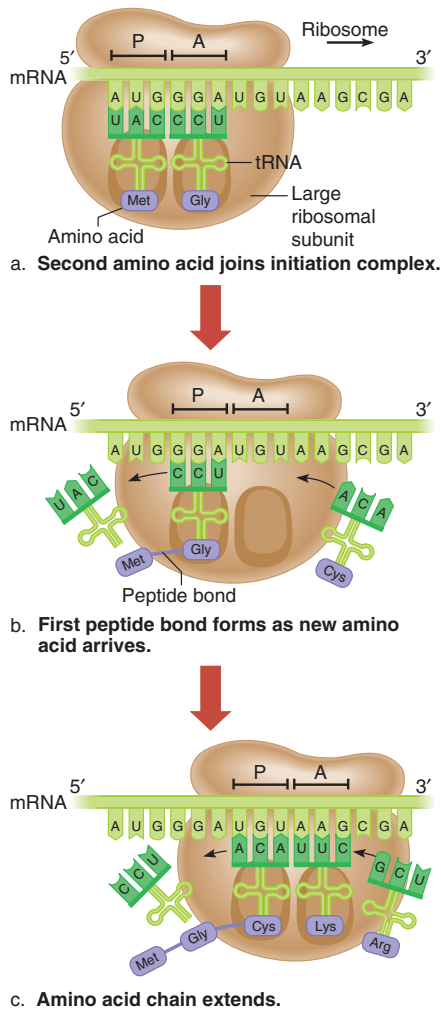


Figure 10.16 Building a polypeptide. (a) A large ribosomal subunit binds to the initiation complex, and a tRNA bearing a second amino acid (glycine, in this example) forms hydrogen bonds between its anticodon and the mRNA's second codon at the A site. The first amino acid, methionine, occupies the P site. (b) The methionine brought in by the first tRNA forms a peptide bond with the amino acid brought in by the second tRNA, and a third tRNA arrives, in this example carrying the amino acid cysteine, at the temporarily vacated A site. (c) A fourth and then fifth amino acid are linked to the growing polypeptide chain. The process continues until a termination codon is reached.

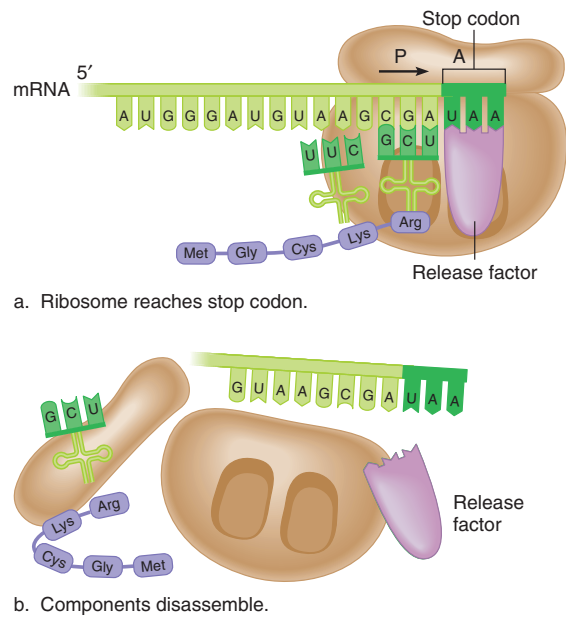


Figure 10.17 Terminating a polypeptide. (a) A protein release factor binds to the stop codon, releasing the completed polypeptide from the tRNA and (b) freeing all of the components of the translation complex.

molecules correspond to it. A protein release factor starts to free the polypeptide. The last tRNA leaves the ribosome, the ribosomal subunits separate and are recycled, and the new polypeptide is released (figure 10.17). Table 10.5 reviews the forms of information encountered in transcription and translation.

Protein synthesis is economical. A cell can produce large amounts of a particular protein from just one or two copies of a gene. A plasma cell in the immune system, for example, manufactures 2,000 identical antibody molecules per second. To mass produce proteins at this rate, RNA, ribosomes, enzymes, and other proteins are continually recycled. In addition, transcription always produces many copies

Table 10.5 Information in Nucleic Acid Molecules

Type of Molecule	Rules and Relationships
DNA coding strand	1 Coding and template strands have complementary DNA bases.
	2 mRNA is complement of DNA template strand, with U for T.
DNA template strand	3 mRNA is same as DNA coding strand, with U for T.
	4 tRNA anticodons are complement of mRNA.
mRNA codons	5 tRNA anticodons are same as DNA template strand, with U for T.
	6 tRNA anticodons are complement of DNA coding strand, with U for T.
tRNA anticodons	7 tRNA translates genetic code, bringing together amino acids specified by DNA coding strand.
Amino acids (protein)	8 Amino acids bond to form a protein.

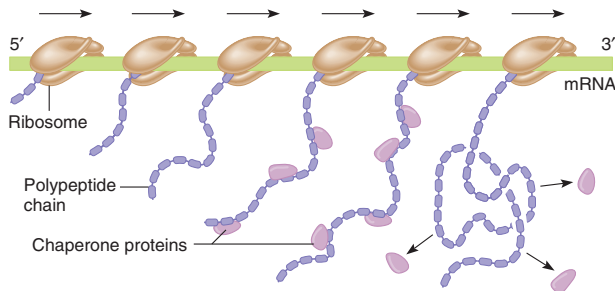


Figure 10.18 Making many copies of a protein. Several ribosomes can simultaneously translate a protein from a single mRNA. These ribosomes hold different-sized polypeptides—the closer to the end of a gene, the longer the polypeptide. Proteins called chaperones help fold the polypeptide.

of a particular mRNA, and each mRNA may bind dozens of ribosomes, as **figure 10.18** shows. As soon as one ribosome has moved far enough along the mRNA to leave space, another ribosome attaches. In this way, many copies of the encoded protein are made from the same mRNA.

As complex as protein synthesis is, linking amino acids is only a first step. The chain must fold in a precise sequence of steps for the protein to assume its three-dimensional form, which is essential for it to function. This may occur in the cytoplasm or on the membranes of the ER, as discussed in the next section.

Some proteins undergo further alterations, called post-translational modifications, before they can function. For example insulin, which is 51 amino acids long, is initially translated as the polypeptide proinsulin, which is 80 amino acids long. Enzymes cut it to 51. Some proteins must have sugars attached for them to become functional, or polypeptides must aggregate.

Key Concepts

1. The genetic code is triplet, nonoverlapping, continuous, universal, and degenerate.
2. As translation begins, mRNA, tRNA with bound amino acids, ribosomes, energy molecules, and protein factors assemble. The mRNA binds to rRNA in a small ribosomal subunit. The first codon attracts a tRNA bearing methionine.
3. The large ribosomal subunit attaches and the tRNA anticodons bind to successive codons. Aligned amino acids form peptide bonds. A polypeptide forms.
4. The ribosome moves down the mRNA to the amino acid chain (the P site) and to where a new tRNA binds (the A site).
5. When the ribosome reaches a “stop” codon, protein synthesis ceases. Components are recycled.

10.4 Processing a Protein

Proteins fold into one or more three-dimensional shapes, or **conformations**. This folding is based on chemistry: attraction and repulsion between atoms of the proteins as well as interactions of proteins with chemicals in the immediate environment. For example, thousands of water molecules surround a growing chain of amino acids. Because some amino acids are attracted to water and some are repelled by it, the water contorts the protein’s shape. Sulfur atoms also affect protein conformation by bridging the two types of amino acids that contain them.

The conformation of a protein is described at several levels (**figure 10.19**). The amino acid sequence of a polypeptide chain is its **primary (1°) structure**. Chemical attractions between amino acids that are close together in the 1° structure fold the polypeptide chain into its **secondary (2°) structure**, which may form loops, coils, barrels, helices, sheets, or other distinctive shapes. The two most common secondary structures are an alpha helix and a beta-pleated sheet. Secondary structures wind into larger **tertiary (3°) structures** as more widely separated amino acids attract or repel in response to water molecules. Finally, proteins consisting of more than one polypeptide form a **quaternary (4°) structure**. Hemoglobin, the blood protein that carries oxygen, has four polypeptide chains (see figure 11.1). The liver protein ferritin has 20 identical polypeptides of 200 amino acids each. In contrast, the muscle protein myoglobin is a single polypeptide chain.

Mutations or polymorphisms may alter the primary structure of a protein if the genetic change is nonsynonymous, which means that it changes the amino acid, just as a non-synonym in the English language is a word that does not have the same meaning as another. In contrast, more than one tertiary or quaternary structure may be possible for a single primary structure because the amino acid chain can fold in different ways. One of the enzymes that is part of the heme pathway depicted in figure 5.6, for example, is normally present as both a very active octamer, which consists of an aggregate of eight molecules of the enzyme, and a low-activity hexamer, which consists of six molecules. In ALAD porphyria, however, there is a greater proportion of hexamers than in the wild type condition. As a result, overall the enzyme is less active.

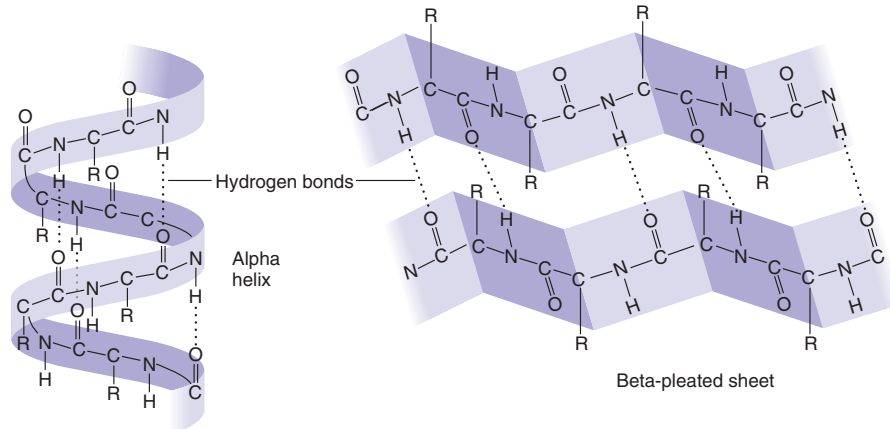
Protein Folding

Protein folding begins within a minute after the amino acid chain winds away from the ribosome. A small protein might contort into its final, functional form in one quick step, taking only a few microseconds. Larger proteins may fold into a series of short-lived intermediates before assuming their final, functional forms.

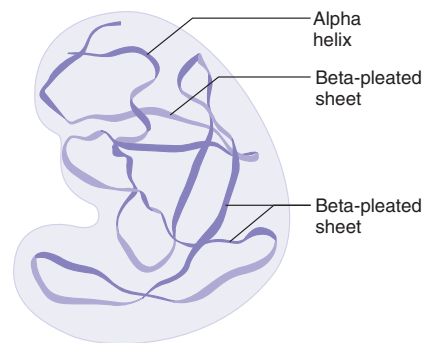
Proteins begin to move toward their destinations as they are being synthesized. In some proteins, part of the start



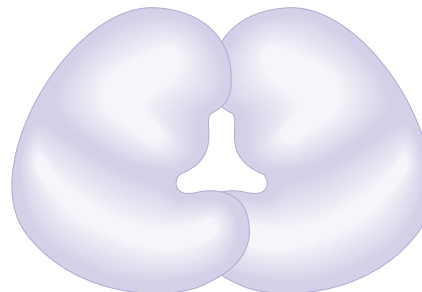
a. **Primary structure**—the sequence of amino acids in a polypeptide chain



b. **Secondary structure**—loops, coils, sheets, or other shapes formed by hydrogen bonds between neighboring carboxyl and amino groups



c. **Tertiary structure**—three-dimensional forms shaped by bonds between R groups, interactions between R groups and water



d. **Quaternary structure**—protein complexes formed by bonds between separate polypeptides

Figure 10.19 Four levels of protein structure. (a) The amino acid sequence of a polypeptide forms the primary structure. Each amino acid has an amino end (NH) and a carboxyl end (COOH), and each of the 20 types of amino acids is distinguished by an R group. (b) Hydrogen bonds between non-R groups create secondary structures such as helices and sheets. The tertiary structure (c) arises when R groups interact, folding the polypeptide in three dimensions and forming a unique shape. (d) If different polypeptide units must interact to be functional, the protein has a quaternary structure.

of the amino acid chain forms a tag of sorts that helps direct the protein in the cell. The first few amino acids in a protein that will be secreted or lodge in a membrane form a “signal sequence” that leads it and the ribosome to which it binds into a pore in the ER membrane. Once in the ER, the protein enters the secretory network (see figure 2.5). Proteins destined for the mitochondria bear a different signal sequence. (Mitochondria manufacture their own proteins but also use many proteins that are encoded in DNA sequences in the nucleus.)

Signal sequences are not found on proteins synthesized on free ribosomes in the cytoplasm. These proteins

may function right where they are made, such as the protein tubules and filaments of the cytoskeleton (see figure 2.10) or enzymes that take part in metabolism. Some proteins travel to and function in the nucleus, such as transcription factors. Proteins destined for the nucleus are synthesized on free ribosomes.

Various proteins assist in this precise folding, whatever the destination. **Chaperone proteins** stabilize partially folded regions in their correct form, and prevent a protein from getting “stuck” in an intermediate form, which would affect its function. Other proteins help new chemical bonds to form

as the final shape arises, and yet others monitor the accuracy of folding. Should a protein misfold, an “unfolded protein response” occurs. Protein synthesis slows or even stops, transcription of genes that encode chaperone proteins and the other folding proteins speeds up, and proper protein folding is usually quickly restored.

Protein Misfolding

If a misfolded protein is made despite these protections, other actions ensue. Misfolded proteins are sent out of the ER back into the cytoplasm, where they are “tagged” with yet another protein, called ubiquitin. A misfolded protein bearing just one ubiquitin tag may straighten and refold correctly, but a protein with more than one tag is taken to another cellular machine called a **proteasome** (figure 10.20). A proteasome is a tunnel-like multi-protein structure. As a protein moves through the opening, it is stretched out, chopped up, and its peptide pieces degraded into amino acids. This is one source of amino acids for building new proteins.

Proteasomes also destroy properly folded proteins that are in excess or no longer needed, perhaps because their job is done. For example, a cell must dismantle excess transcription factors, or the genes that they control may remain activated or repressed for too long. Proteasomes also control the abundance of cyclin proteins, which must be present in certain numbers to maintain the cell cycle. They also dismantle proteins from pathogens, such as viruses.

Most misfolded proteins are the result of errors in any of the steps of protein synthesis and processing. The primary structure (amino acid sequence) may be wild type or a functional variant, but the process goes awry as the proteins fold. Or, a mutation may alter the primary structure in a way that affects attractions and repulsions between parts of

the protein. A clear example of protein misfolding in a single-gene disorder is sickle cell disease (see figure 12.2). Instead of the normal globular conformation of hemoglobin (see figure 11.1), the protein forms sheets that bend the red blood cells that contain them out of shape. **Table 10.6** lists some disorders that, in some cases, involve protein misfolding.

Some mutations that cause cystic fibrosis prevent CFTR protein from folding and anchoring in the plasma membrane, where it normally controls the flow of chloride ions. Instead, the misfolded protein builds up in the cell.

In several disorders that affect the brain, misfolded proteins aggregate, forming masses that clog the proteasomes and block them from processing any malformed proteins. Different proteins are affected in different disorders. In Huntington disease, for example, extra glutamines in the protein huntingtin cause it to obstruct proteasomes. Misfolded proteins that clog proteasomes also form in the disorders listed in Table 10.6, but it isn't always clear whether the accumulated proteins cause the disease or are a response to it. Some of these disorders are discussed further in chapter 12.

Understanding how protein misfolding causes diseases can lead to development of new treatments, or recognition that existing drugs may help. This is the case for phenylketonuria (PKU), discussed briefly in chapter 2.

PKU

The story of PKU began in Oslo, Norway, in 1934, when an observant mother of two children with extreme mental retardation noticed that the children's soiled diapers had an odd, musty odor. She mentioned this to Ivar Fölling, a relative who was a physician and a biochemist. Intrigued, Fölling analyzed the children's urine and found excess phenylalanine, an amino acid usually scant in urine because an enzyme, phenylalanine

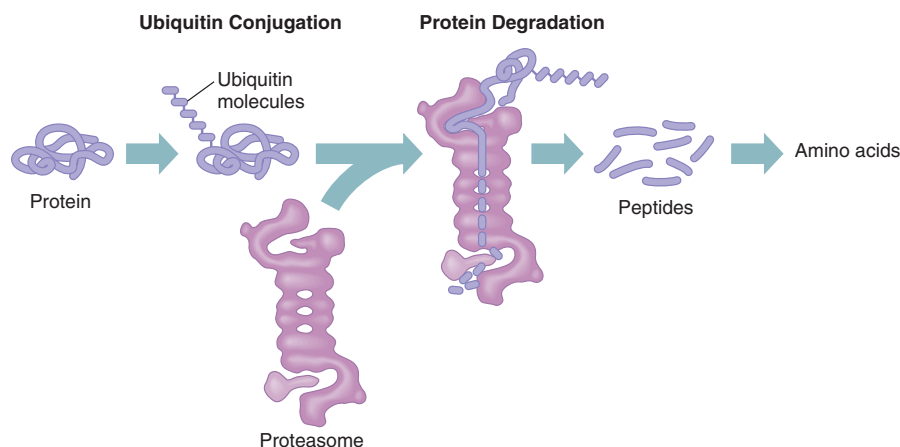


Figure 10.20 Proteasomes provide quality control. Ubiquitin binds to a misfolded protein and escorts it to a proteasome. The proteasome, which is composed of several proteins, encases the misfolded protein, straightening and dismantling it.

Table 10.6 Disorders Associated with Protein Misfolding

Disease	Misfolded Protein	MIM (protein)
Alzheimer disease	Amyloid beta precursor protein	104760
Familial amyotrophic lateral sclerosis	Superoxide dismutase	147450
Huntington disease	Huntingtin	143100
Parkinson disease	Alpha synuclein	163890
Lewy body dementia	Alpha synuclein	
PKU	Phenylalanine hydroxylase	261600
Prion disorders	Prion protein	176640

(All but Huntington disease are genetically heterogeneous; that is, abnormalities in different proteins cause similar syndromes.)

hydroxylase (PAH), breaks it down. The children's cells lacked enzyme activity because the children had inherited PKU from their carrier parents. The build-up of phenylalanine affected their brains.

In 1963, a dietary treatment was devised that supplies protein in a synthetic food that is very low in phenylalanine. Begun at birth, the diet prevents symptoms. However, it is difficult to follow, and older children and teens often "cheat." When they eat protein, symptoms can surface, such as slowed

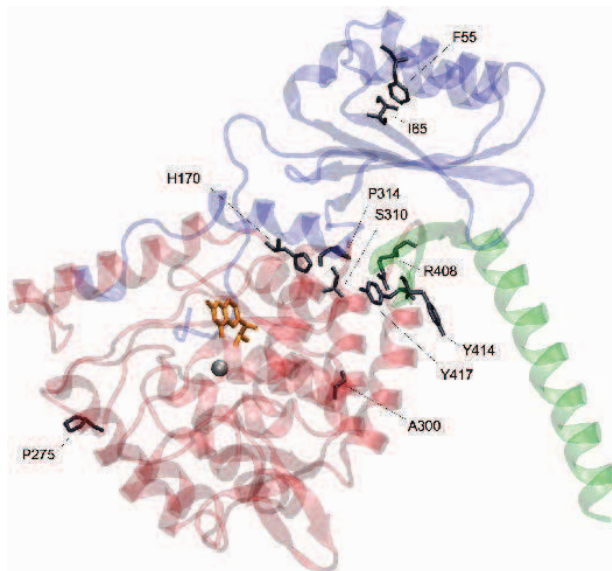


Figure 10.21 Absence of activity of an enzyme (phenylalanine hydroxylase) causes PKU. Mental retardation results unless the person follows a diet that limits the amino acid phenylalanine, which the enzyme normally breaks down. In the illustration, the superimposed black regions are the parts where mutations affect the protein.

reaction times, memory deficits, anxiety, depression, and irritability. In 2002, another treatment was found to help people with mild cases of PKU—the vitamin derivative tetrahydrobiopterin, or BH_4 . This small molecule usually nestles into the four proteins that assemble to form PAH. In patients with mild cases of PKU, taking BH_4 increases enzyme levels, presumably restoring function in some of the enzyme. Researchers then discovered that even some patients with more serious PKU were helped with a drug version of BH_4 . What was happening? Protein misfolding explains the success of the treatment.

Each of the four subunits of PAH, the enzyme thought to be missing in severe cases of PKU, may not actually be missing. Instead, enzyme molecules are so misfolded that they cannot work. When a person has a mutation, shown in the black outlines in **figure 10.21**, localized misfolding occurs. Because of the tertiary structure, the disturbance spreads to amino acids that are actually far away in the primary sequence, but close by when the protein is folded. Misfolding spreads until the entire, four-protein PAH enzyme can no longer function. The peculiarities of the tertiary structure of this particular molecule enable it to assume any of several conformations, making it prone to misfolding.

Prion Diseases

Several disorders that affect the brain reflect alternate folding of a single glycoprotein called a prion, pronounced *pree-on* (*PRNP*, MIM 176640). Like the enzyme behind PKU, prion protein (PrP) can also fold into any of several conformations—probably at least eight. What distinguishes the several prion disorders, however, is that one conformation is “infectious,” causing other prions that it contacts to also assume the infectious form. Another distinguishing feature of the prion diseases is that the infectious forms can come from even the wild type primary structure.

Prion diseases are called transmissible spongiform encephalopathies (TSEs) (**figure 10.22** and Reading 10.1). In the 85 species in which TSEs are known, the brains become riddled with holes, resembling a sponge. Nerve cells die, neuroglial cells overgrow, and death occurs within 18 months of the first symptom. The location of the damage determines the specific symptoms, but most TSEs cause extreme weight loss, poor coordination, and dementia. One disease, fatal familial insomnia, causes death in months from inability to sleep. It also causes tremor, fever, sweats, muscle aches, joint pain, seizures, and a dreamlike trance.

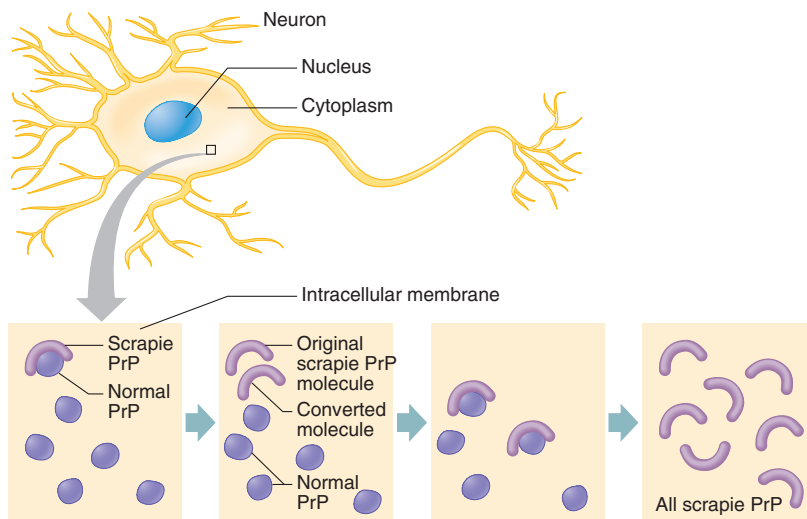


Figure 10.22 Prions change shape. A prion disease may begin when a single scrapie prion protein (PrP) contacts a normal PrP and changes it into the scrapie conformation. As the change spreads, disease results, usually with accumulated scrapie prion proteins clogging brain tissue.

TSEs were discovered in sheep, which develop a disease called scrapie when they eat prion-infected brains from other sheep. Ten percent of TSE cases in humans are inherited, such as Creutzfeldt-Jakob disease and Gerstmann-Straussler disease. Other cases are acquired from ingesting prions, or sporadic, occurring without a family history and with no apparent source. In acquired TSEs, exposure to prions in the infectious conformation triggers conversion of the person's own normal prions into the disease-causing conformation.

The “rules” by which DNA sequences specify protein conformations are still not well understood, even as we routinely decipher and describe the sequences of entire genomes. In fact, sequencing the human genome has revealed that encoding protein is only one function of DNA. Even though the central dogma remains true—genes are transcribed into mRNA, which, in turn, is translated into protein—many transcripts are not represented in the proteins encoded in the genome. What do they do? The next chapter offers some ideas. Amidst all the new questions that deciphering the human genome sequence inspired is one inarguable fact: The straightforward, linear relationship between gene and protein that emerged from the experiments of the 1960s was merely an opening chapter to the story of how cells survive, specialize, and interact as they build bodies.

Key Concepts

1. Protein folding begins as translation proceeds, with enzymes and chaperone proteins assisting.
2. Misfolded proteins are tagged with ubiquitin and sent through a proteasome for dismantling.
3. A protein can fold in more than one way. Some conformations cause disease.

Reading 10.1

Considering Kuru

A few rare and strange disorders are caused by *prions*—proteins that can infect.

The first prion disease of humans to be recognized was kuru, which struck the Foré people in a remote mountainous area of Papua New Guinea (figure 1). In the Foré language, *kuru* means to shake. The disease began with wobbling legs, quickly followed by trembling hands and fingers, and then body-wide shaking. An odd sign was uncontrollable laughter, leading to the nickname “laughing disease.” Speech slurred and faded, thinking slowed, and after several months, the person could no longer walk or eat. Death typically came within a year. One in 10 of the 35,000 tribe members had kuru.

The fact that only women and young children developed kuru at first suggested that the disease might be inherited, but D. Carleton Gajdusek, a physician who spent much of his lifetime studying the Foré, learned that the preparation of human brains for a cannibalism ritual probably passed on the infectious prions. People ate their relatives killed in battle to honor them. After the ritual was banned in 1959, the disease gradually disappeared. Gajdusek vividly described the Foré preparation of human brains at a time when he thought the cause was viral:

Children participated in both the butchery and the handling of cooked meat, rubbing their soiled hands in their armpits

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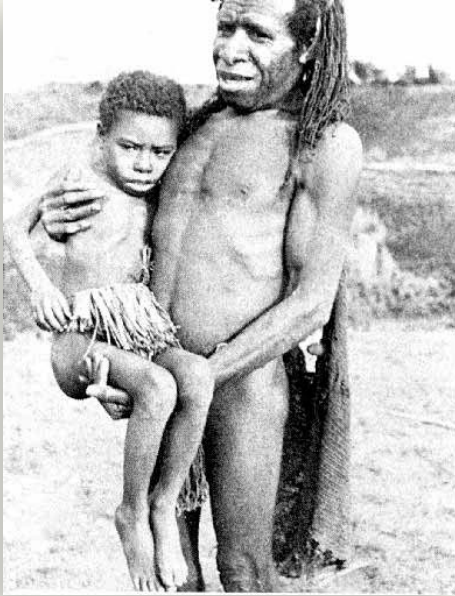


Figure 1 Kuru. Kuru is a prion disease that affected the Foré people of New Guinea until they gave up a cannibalism ritual that spread an infectious form of prion protein.

or hair, and elsewhere on their bodies. They rarely or never washed. Infection with the kuru virus was most probably through the cuts and abrasions of the skin or from nose picking, eye rubbing, or mucosal injury.

Although kuru vanished, other prion diseases surfaced (**table 1**). In the 1970s and 1980s, several people acquired Creutzfeldt-Jakob disease (CJD). This time, the route of transmission was either through corneal transplants, in which infectious prions entered the brain through the optic nerve, or from human growth hormone taken from cadavers and used to treat short stature in children. The most familiar prion disease is “mad cow disease” and the variant CJD it has caused in more than 120 people in the United Kingdom since 1995. People likely acquired the infectious prions by eating infected beef.

Table 1 Prion Disorders

Disorder	MIM #
Creutzfeldt-Jakob disease	123400
Fatal familial insomnia	600072
Gerstmann-Straussler Disease	137440

Summary

10.1 From DNA to Protein

1. A small part of the genome encodes protein. Much of the rest controls protein synthesis.

10.2 Transcription

2. Some DNA is **transcribed** into RNA, which is then **translated** into protein.
3. RNA is transcribed from the **template strand** of DNA. The other DNA strand is called the **coding strand**.
4. RNA is a single-stranded nucleic acid similar to DNA but containing uracil and ribose rather than thymine and deoxyribose.
5. **Messenger RNA** (mRNA) carries a protein-encoding gene’s information. **Ribosomal RNA** (rRNA) associates with certain proteins to form ribosomes, which physically support protein synthesis. **Transfer RNA** (tRNA) is cloverleaf-shaped, with a three-base **anticodon** that is complementary to mRNA on one end and bonds to a particular amino acid on the other end.
6. Operons control gene expression in bacteria. In more complex organisms, **transcription factors** regulate which genes are transcribed in a particular cell type.
7. Transcription begins when transcription factors help **RNA polymerase** (RNAP) bind to a gene’s **promoter**. RNAP then

adds RNA nucleotides to a growing chain, in a sequence complementary to the DNA template strand.

8. After a gene is transcribed, the mRNA receives a “cap” of modified nucleotides at the 5’ end and a poly A tail at the 3’ end.
9. Many genes do not encode information in a continuous manner. After transcription, **exons** are translated into protein and **introns** are removed. Introns may outnumber and outsize exons. **Alternate splicing** increases protein diversity.

10.3 Translation of a Protein

10. Each three consecutive mRNA bases form a **codon** that specifies a particular amino acid. The **genetic code** is the correspondence between each codon and the amino acid it specifies. Of the 64 different possible codons, 60 specify amino acids, one specifies the amino acid methionine and “start,” and three signal “stop.” Because 61 codons specify the 20 amino acids, more than one type of codon may encode a single amino acid. The genetic code is nonoverlapping, triplet, universal, and degenerate.
11. In the 1960s, researchers used logic and clever experiments using synthetic RNAs to decipher the genetic code.
12. Translation requires tRNA, ribosomes, energy-storage molecules, enzymes, and protein factors. An initiation complex forms when mRNA, a small ribosomal subunit, and a tRNA carrying methionine join. The amino acid chain

elongates when a large ribosomal subunit joins the small one. Next, a second tRNA binds by its anticodon to the next mRNA codon, and its amino acid bonds with the first amino acid. Transfer RNAs add more amino acids, forming a polypeptide. The ribosome moves down the mRNA as the chain grows. The P site bears the amino acid chain, and the A site holds the newest tRNA. When the ribosome reaches a “stop” codon, it falls apart into its two subunits and is released. The new polypeptide breaks free.

- After translation, some polypeptides are cleaved, have sugars added, or aggregate. The cell uses or secretes the protein.

10.4 Processing a Protein

- A protein must fold into a particular **conformation** to be active and functional.

- A protein’s **primary structure** is its amino acid sequence. Its **secondary structure** forms as amino acids close in the primary structure attract one another. **Tertiary structure** appears as more widely separated amino acids attract or repel in response to water molecules. **Quaternary structure** forms when a protein consists of more than one polypeptide.
- Chaperone proteins** help conformation arise. Other proteins help new bonds form and oversee folding accuracy.
- Ubiquitin attaches to misfolded proteins and escorts them to **proteasomes** for dismantling. Protein misfolding is associated with certain diseases.
- Some proteins can fold into several conformations, some of which can cause disease.
- At least one conformation of prion protein is infectious, causing transmissible spongiform encephalopathies.

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Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

- Explain how complementary base pairing is responsible for
 - the structure of the DNA double helix.
 - DNA replication.
 - transcription of RNA from DNA.
 - the attachment of mRNA to a ribosome.
 - codon/anticodon pairing.
 - tRNA conformation.
- A retrovirus has RNA as its genetic material. When it infects a cell, it uses enzymes to copy its RNA into DNA, which then integrates into the host cell’s chromosome. Is this flow of genetic information consistent with the central dogma? Why or why not?
- What are the functions of these proteins?
 - RNA polymerase
 - ubiquitin
 - a chaperone protein
 - a transcription factor
- Explain where a hydrogen bond forms and where a peptide bond forms.
- List the differences between RNA and DNA.
- Where in a cell do DNA replication, transcription, and translation occur?
- How does transcription control cell specialization?
- How can the same mRNA codon be at an A site on a ribosome at one time, but at a P site at another time?
- Describe the events of transcription initiation.
- List the three major types of RNA and their functions.
- Describe three ways RNA is altered after it is transcribed.
- What are the components of a ribosome?
- Why would an overlapping genetic code be restrictive?
- Why would two-nucleotide codons be insufficient to encode the number of amino acids in biological proteins?
- How are the processes of transcription and translation economical?
- Explain how protein misfolding conditions and illnesses that result from abnormal transcription factors might each produce many different symptoms.
- What factors determine how a protein folds into its characteristic conformation?
- How do a protein’s primary, secondary, and tertiary structures affect conformation? Which is the most important determinant of conformation?

Applied Questions

- List the RNA sequence transcribed from the DNA template sequence TTACTTGGCTTGAGATC.
- Reconstruct the corresponding DNA template sequence from the partial mRNA sequence GCUAUCUGUCAUAAAAGAGGA.

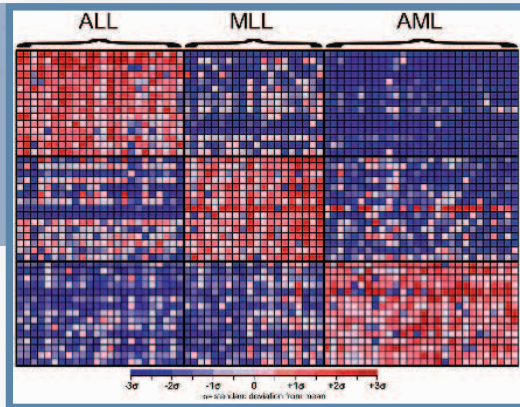
- List three different mRNA sequences that could encode the amino acid sequence
histidine-alanine-arginine-serine-leucine-valine-cysteine.
- Write a DNA sequence that would encode the amino acid sequence valine-tryptophan-lysine-proline-phenylalanine-threonine.
- In the film *Jurassic Park*, which is about cloned dinosaurs, a cartoon character named Mr. DNA talks about the billions of genetic codes in DNA. Why is this statement incorrect?
- Titin is a muscle protein named for its size—its gene has the largest known coding sequence of 80,781 DNA bases. How many amino acids long is it?
- An extraterrestrial life form has a triplet genetic code with five different bases. How many different amino acids can this code specify, assuming no degeneracy?
- In malignant hyperthermia, a person develops a life-threateningly high fever after taking certain types of anesthetic drugs. In one family, mutation deletes three contiguous bases in exon 44. How many amino acids are missing from the protein?
- The protein that serves as a receptor that allows insulin to enter cells has a different number of amino acids in a fetus and in an adult. Explain how this may happen.

Web Activities

- Go to <http://www.mcb.harvard.edu/BioLinks/gencode.html>. Scroll down to the lists of “noncanonical” codes in organisms other than humans. (*Noncanonical* means it differs from the universal genetic code.) Find three examples of deviations from the universal code, and list what the codon-amino acid assignment is in most organisms. (Replace the T’s on the website with the U’s to correspond to the genetic code chart in the textbook.)
- Use the Web to find out how the ubiquitin-proteasome system is overtaxed or disabled in a neurodegenerative disease such as Alzheimer disease, Parkinson disease, Huntington disease, amyotrophic lateral sclerosis, or Lewy body dementia. (Find websites for these disorders and discuss how the mechanism involves proteasomes.)

Case Studies and Research Results

- Five patients meet at a clinic for families in which several members have early-onset Parkinson disease. This condition causes rigidity, tremors, and other motor symptoms. Only 2 percent of cases of Parkinson disease are inherited. The five patients all have mutations in a gene that encodes the protein parkin, which has 12 exons. For each patient, indicate whether the mutation shortens, lengthens, or does not change the size of the protein.
 - Manny Filipo’s *parkin* gene is missing exon 3.
 - Frank Myer’s *parkin* gene has a duplication in intron 4.
 - Theresa Ruzi’s *parkin* gene lacks six contiguous nucleotides in exon 1.
 - Elyse Fitzsimmon’s *parkin* gene has an altered splice site between exon 8 and intron 8.
 - Scott Shapiro’s *parkin* gene is deleted.
- A research project called ENCODE (ENCyclopedia of DNA Elements) took 1 percent of the human genome and cataloged 487 genes and 2,608 mRNA transcripts. Why isn’t the number the same?



Colors represent different levels of gene expression. These leukemias—ALL, MLL, and AML—differ in gene expression patterns. The vertical columns of squares represent DNA from tumor samples, and the horizontal rows compare the activities of particular genes. Red tones indicate higher-than-normal expression and blue tones show lower-than-normal expression. The different patterns indicate distinct cancers, although the cells look alike under a microscope.

CHAPTER

11

Gene Expression and Epigenetics

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Uncloaking a Cancer

Ten years ago, Tim and Donna met at the medical center where each went every few weeks to receive chemotherapy. Tim was 4, Donna still a baby, but he enjoyed sharing picture books with her. Both had been diagnosed with acute lymphoblastic leukemia (ALL) following several weeks of fatigue, fever, and bruising. Their cancerous white blood cells were dividing too frequently, crowding out red blood cells and platelets in their bone marrow. Both children received the same chemotherapy. The only bright spot in Tim's hospital visits was seeing his little friend.

Then one day Donna was not there. She had died. The chemotherapy had helped Tim, as it does 90 percent of children with ALL, and he recovered. But Donna had only gotten sicker.

Researchers at the Dana Farber Cancer Institute wondered why some young children with ALL do not respond to the same treatment that helps many others. So they examined what the cancer cells *do*, rather than how they appear. They used DNA microarrays ("chips") to compare the expression of 12,000 genes in cancer cells from ALL patients who were helped by the standard therapy and patients who weren't. Compared to patients like Tim who responded, the cancerous cells of patients like Donna made too little of 1,000 of the encoded proteins and too many of 200 others. The children who didn't respond had a different type of disease, even though they had the same symptoms and their cancer cells looked alike. At the level of gene expression—the proteins manufactured in the cancer cells—the diseases were distinctive (see figure).

The 10 percent of children who do not respond to ALL treatment actually have MLL—mixed-lineage leukemia. Other drugs can help them.

11.1 Gene Expression Through Time and Tissue

A genome is like an orchestra. Just as not all of the instruments play with the same intensity at every moment, not all genes are expressed continually at the same levels. Before the field of genomics began in the 1990s, the study of genetics proceeded one gene at a time, like hearing the separate contributions of a violin, a viola, and a flute. Many genetic investigations today, in contrast, track the crescendos of gene activity that parallel events in an organism's life. This new view has introduced the element of time to genetic analysis. Unlike the gene maps of old, which ordered genes on chromosomes, new types of maps reveal the timing of gene expression in unfolding programs of development and response to the environment.

The discoveries of the 1950s and 1960s on DNA structure and function answered some questions about the control of gene expression while raising many more. How does a bone cell “know” to transcribe the genes that control the synthesis of collagen and not to transcribe genes that specify muscle proteins? What causes the proportions of blood cell types to shift into leukemia? How do chemical groups “know” to shield DNA from transcription in one circumstance, yet expose it in others?

Changes to the chemical groups that associate with DNA profoundly affect which parts of the genome are accessible to transcription factors and under which conditions. Such changes to the molecules that bind to DNA that are transmitted to daughter cells when the cell divides are termed **epigenetic**—literally, “outside the gene.”

Specific classes of proteins and RNA molecules carry out epigenetic changes, and much of the genome encodes these modifiers of gene expression. Although research is currently focusing on identifying epigenetic changes in different cell types under different conditions, the idea of modifying gene expression is not new. British embryologist C. H. Waddington wrote in 1939 of “the causal interactions between genes and their products which bring the phenotype into being.” Epigenetic changes do not alter the DNA base sequence, although they are passed from one cell generation to the next.

This chapter extends the discussion of DNA structure and function from the encoding of information to how that information is accessed. We begin with three examples at the molecular, tissue, and organ levels: (1) hemoglobin switching during development; (2) the composition of blood plasma, and; (3) specialization of the two major parts of the pancreas.

Globin Chain Switching

The globin proteins that transport oxygen in the blood vividly illustrate control of gene expression. The molecule's changing composition through development was discovered half a century ago.

A hemoglobin molecule in the blood of an adult has four polypeptide chains, each wound into a globular conformation (**figure 11.1**). Two of the chains are

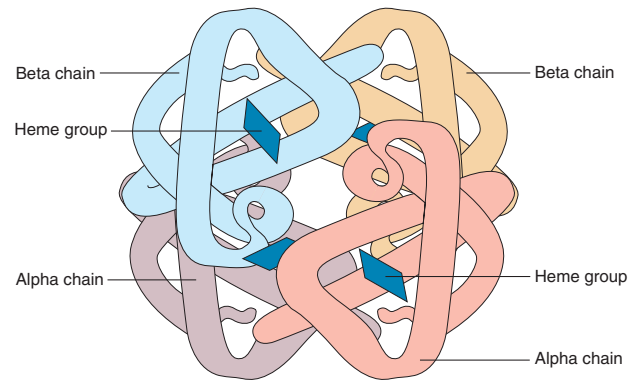


Figure 11.1 The structure of hemoglobin. A hemoglobin molecule is made up of two globular protein chains from the beta (β) globin group and two from the alpha (α) globin group. Each globin surrounds an iron-containing chemical group called a heme.

146 amino acids long and are called “beta” (β). The other two chains are 141 amino acids long and are termed “alpha” (α). The genes for beta subunits are clustered on chromosome 11, and the alpha genes are grouped on chromosome 16.

The subunits of the hemoglobin molecule are replaced as the oxygen concentration in the body changes, which in turn depends upon whether oxygen arrives to an embryo or fetus through the placenta or to a newborn's lungs from breathing. The chemical basis for this “globin chain switching” is that different polypeptide subunits attract oxygen molecules to different degrees. Parts of the globin gene clusters oversee the changes in the molecule's composition and assembly.

The subunit makeup of the hemoglobin molecule differs in the embryo, fetus, and adult (**figure 11.2**). In the embryo,

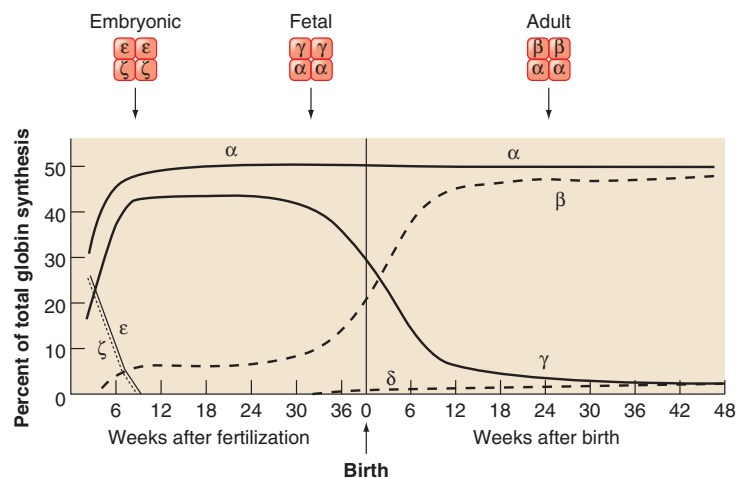


Figure 11.2 Globin chain switching. The subunit composition of human hemoglobin changes as the concentration of oxygen in the environment changes. With the switch from the placenta to the newborn's lungs to obtain oxygen, beta (β) globin begins to replace gamma (γ) globin.

as the placenta forms, hemoglobin consists first of two epsilon (ϵ) chains, which are in the beta globin group, and two zeta (ζ) chains, which are in the alpha globin group. About 4 percent of the hemoglobin in the embryo includes beta chains. This percentage gradually increases.

As the embryo develops into a fetus, the epsilon and zeta chains decrease in number, as gamma (γ) and alpha chains accumulate. Hemoglobin consisting of two gamma and two alpha chains is called fetal hemoglobin. The gamma globin subunits bind very strongly to oxygen released from maternal red blood cells into the placenta, so that fetal blood carries 20 to 30 percent more oxygen than an adult's blood. As the fetus matures, beta chains gradually replace the gamma chains. At birth, however, the hemoglobin is not fully of the adult type—fetal hemoglobin (two gamma and two alpha chains) comprises from 50 to 85 percent of the blood. By four months of age, the proportion drops to 10 to 15 percent, and by four years, it is less than 1 percent.

Building Tissues and Organs

The globin chains affect one type of molecule, hemoglobin. Changing gene expression and the resulting production of proteins can also be observed on a larger scale, such as the numbers of different types and amounts of proteins in particular tissues. For example, blood plasma, the liquid portion of blood, contains about 40,000 different types of proteins. Ten types of proteins account for 90 percent of all the plasma protein molecules, and nearly half of those are one type, albumin. Many thousands of types of proteins make up the rest, and are present in very small amounts. If conditions change, such as a person developing an infection or allergic reaction, the protein profile of the plasma can change dramatically. This ability of the tissue to adapt to a changing environment is possible because of changes in gene expression—that is, how much of each protein is made.

Blood is a structurally simple tissue that is easy to obtain and study. A solid gland or organ, constructed from specialized cells and tissues, is much more complex. Its solid organization must be maintained throughout a lifetime of growth, repair, and changing external conditions.

Stem cell biology is shedding light on how genes are turned on and off during the development of an organ or gland. Researchers isolate individual stem cells and then

see which combinations of growth factors, hormones, and other biochemicals must be added to steer development toward a particular cell type.

Consider the pancreas. It is a dual gland, with two types of cell clusters. The exocrine part releases digestive enzymes into ducts, whereas the endocrine part secretes polypeptide hormones that control nutrient use directly into the bloodstream. The endocrine cell clusters are called pancreatic islets.

The complexity of the pancreas unfolds in the embryo, when ducts form. Within duct walls reside rare stem cells and progenitor cells (see figure 2.22). Recall that a stem cell is an unspecialized cell that divides to yield another stem cell (self-renewal), and a progenitor cell that is partially specialized. When a transcription factor called *pdx-1* is activated, it in turn controls expression of other genes in a way that stimulates some of the progenitor cells to divide. Some of the progenitor cells give rise to daughter cells that follow an exocrine pathway; they are destined to produce digestive enzymes (**figure 11.3**). Other progenitor cells respond to different signals and divide to yield daughters that follow the endocrine pathway. The most familiar pancreatic hormone is

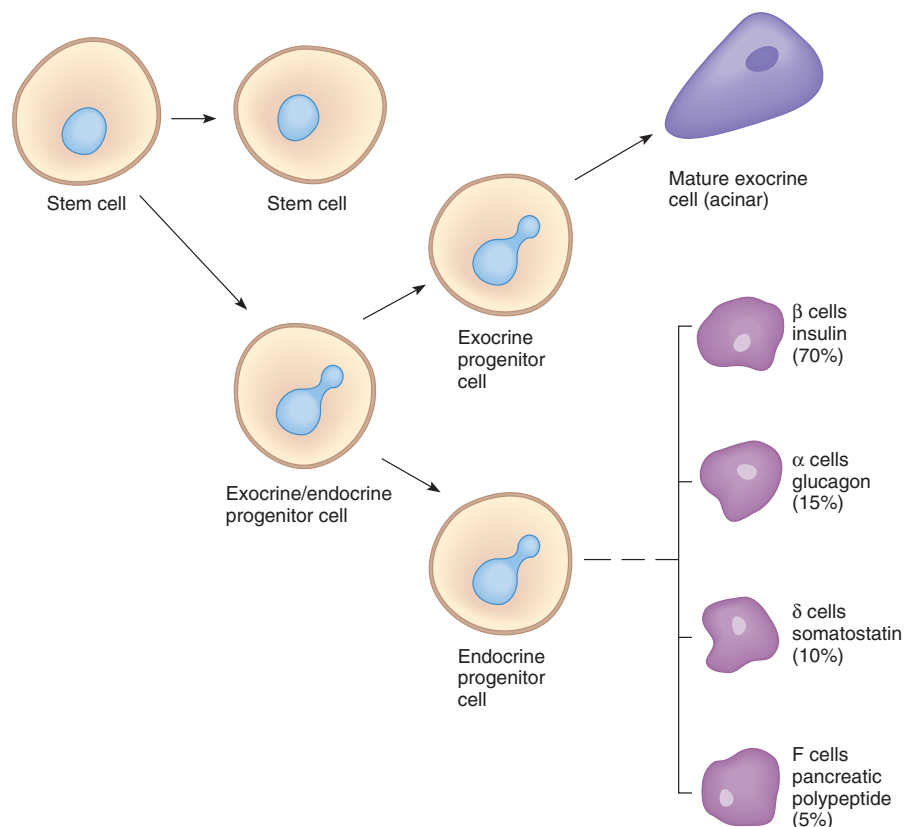


Figure 11.3 Building a pancreas. A single type of stem cell theoretically gives rise to an exocrine/endocrine progenitor cell that in turn divides to yield more restricted progenitor cells that give rise to both mature exocrine and endocrine cells. The endocrine progenitor cell in turn divides to give rise to cells that are specialized to produce particular hormones.

insulin—its absence (or the inability of cells to recognize it) causes diabetes mellitus.

Researchers can observe the specialization of pancreas cells by taking individual progenitor cells from human pancreas ducts or by deriving them from reprogrammed somatic cells (see figure 2.24). Then, by supplying specific growth factors at particular times, the progenitor cells give rise to clusters that look and function like pancreatic islets. When exposed to glucose, the cells secrete insulin! If pancreatic stem cells can be isolated and cultured, it might be possible to coax a person with diabetes to produce new and functional pancreatic beta cells.

Proteomics

A more complete portrait of gene expression emerges through **proteomics**, which considers all proteins made in a cell, tissue, gland, organ, or entire body. **Figure 11.4** depicts a global way of comparing the relative contributions of major categories of proteins from conception through birth and from conception through old age.

The differences in proteins produced at different times make sense. For example, transcription factors are more abundant before birth because of the extensive cell

differentiation of this period. During the prenatal period, enzymes are less abundant, perhaps because the fetus receives some enzymes through the placenta. Immunoglobulins appear as a category after birth as the immune system begins to function.

Another way to look at the proteome is by specific functions, which has led to the creation of various “ome” words. Genes whose encoded proteins control lipid synthesis, for example, constitute the “lipidome,” and those that monitor carbohydrate production and use form the “glycome.” “Omics” designations are helpful in sorting out the thousands of proteins a human cell can manufacture. However, identifying proteins is only a first step. The next hurdle is to determine how proteins with related functions interact—forming “interactomes.”

Gene expression profiles for different cell types under various conditions can provide valuable medical information and are the basis for many new tests that diagnose disease or monitor response to treatment. For example, 55 genes are overexpressed and 480 underexpressed in cells of a prostate cancer that has a very high likelihood of spreading—but not in a prostate cancer that will not spread. A test based on such findings assists physicians in deciding which patients can benefit most from further treatment.

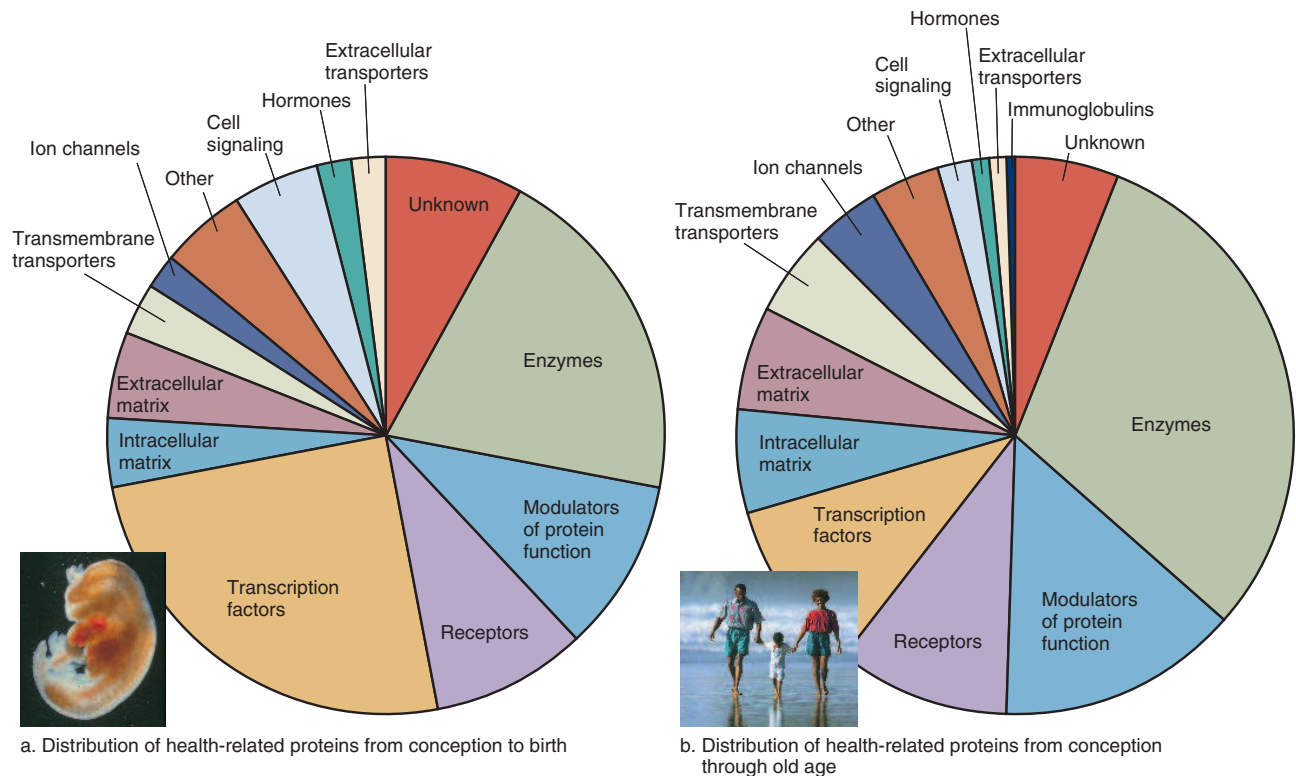


Figure 11.4 Proteomics meets medicine. We can categorize genes by their protein products, and chart the relative abundance of each class at different stages of development. The pie chart in (a) considers 13 categories of proteins that when abnormal or missing cause disease, and their relative abundance from conception to birth. The pie chart in (b) displays the same protein categories from conception to old age, plus one activated after birth, the immunoglobulins.

Key Concepts

1. Gene expression patterns change over time and in different cell types.
2. The subunit composition of hemoglobin changes in the embryo, fetus, and after birth.
3. As a pancreas forms, progenitor cells diverge from shared stem cells and their daughters specialize.
4. Proteomics tracks all of the proteins in a cell, tissue, organ, or organism under specific conditions.

11.2 Control of Gene Expression

Purchasing a complex appliance or device that requires assembly but does not come with clear instructions can be very frustrating. Fortunately, this is not true of the human genome. Embedded in the blueprints to build a human body are several types of instructions for its use.

A protein-encoding gene contains some controls over its own expression level. One is the promoter sequence. Recall from chapter 10 that the promoter is part of the 3' end of a gene where RNA polymerase and transcription factors bind, marking the start point of transcription. Variations in the promoter sequence of a gene can affect how quickly the encoded protein is synthesized. For example, a form of early-onset Alzheimer disease is caused by a mutation in the promoter for the gene that encodes amyloid precursor protein (see Reading 4.1). In people who have the mutation, the sticky protein characteristic of the disease accumulates in the brain twice as fast as normal. A second way that expression of a gene can exceed normal pace is if a person has more than one copy of it.

Much of the control of gene expression occurs in two general processes. In the first, **chromatin remodeling**, interactions of the histone proteins associated with DNA with other chemical groups expose some sections of DNA to transcription factors and shield other sections, blocking their expression. Later in the protein production process, small RNAs called, appropriately, **microRNAs**, bind the 3' ends of certain mRNAs, preventing their translation into protein. Overall, these two processes determine the ebb and flow of different proteins, enabling cells to adapt to changing conditions.

Chromatin Remodeling

Recall that DNA associates with proteins and RNA to form chromatin, as figure 9.13 shows. For many years, biologists thought that the histone proteins that wind long DNA molecules into nucleosomes were little more than tiny spools. However, histones do much more. They play a major role in exposing DNA when and where it is to be transcribed, and shielding it when it is to be silenced. To do this, enzymes add or delete small organic chemical groups to histones in a process called chromatin remodeling. The resulting patterns of chemical groups control the effect of histones on their associated protein-encoding genes.

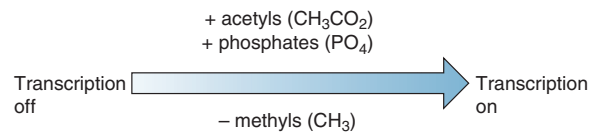


Figure 11.5 Chromatin remodeling. Chromatin remodeling adds or removes certain organic chemical groups to or from histones. The pattern of binding controls whether the DNA wrapped around the histones is transcribed or not.

The three major types of small molecules that bind to histones are acetyl groups, methyl groups, and phosphate groups (figure 11.5). The key to the role histones play in controlling gene expression lies in acetyl groups (CH_3CO_2). They bind to very specific sites on certain histones, particularly to the amino acid lysine.

Figure 11.6 shows how acetyl binding can subtly shift histone interactions in a way that eases transcription. A series of proteins moves the histone complex away from the TATA box, exposing it enough for RNA polymerase to bind and transcription to begin (see figure 10.7). First, a group of proteins called an enhanceosome attracts the enzyme (acetylase) that adds acetyl groups to specific lysines on specific histones. Adding acetyl groups to these lysines neutralizes their positive charge. Because DNA carries a negative charge, this change moves the histones away from the DNA, enabling transcription factors to enter, bind, and begin transcription. Enzymes called deacetylases remove acetyl groups, which shuts off gene expression.

Methyl groups (CH_3) are also added to or taken away from histones. When CH_3 binds to a specific amino acid in a specific histone type, a protein is attracted that shuts the DNA off. As CH_3 groups are added, methylation spreads from the tail of one histone to the adjacent histone, propagating the gene silencing. Methyl groups also control gene expression by binding to cytosines at about 10,000 places in the genome.

The modified state of chromatin can be passed on when DNA replicates. These changes in gene expression are heritable from cell generation to cell generation, but they do not alter the DNA base sequence—that is, they are epigenetic. Addition and removal of acetyl groups, methyl groups, or phosphate groups are examples of epigenetic changes. Effects of methylation can sometimes be seen when MZ (identical) twins inherit the same disease-causing genotype, but only one twin is sick. The reason for the discordance may be different patterns of methylation of the gene.

Enzymes that add or delete acetyl, methyl, and phosphate groups must be in a balance that controls which genes are expressed and which are silenced. Upset this balance, and disease can result. In the mixed lineage leukemia described in the chapter opener, for example, a single abnormal protein binds to more than 150 genes and alters their associated chromatin. Among these genes are several that normally stimulate frequent cell division in the stem cells that give rise to blood cells. In the leukemia, these overexpressed genes send the affected white blood cells back in developmental time, to a state in which the rapid cell division causes cancer. One limitation to altering chromatin remodeling to treat inherited disease is that this action could affect the expression of many genes—not just the one implicated in the disease. Table 11.1 lists disorders that result from abnormal chromatin remodeling.

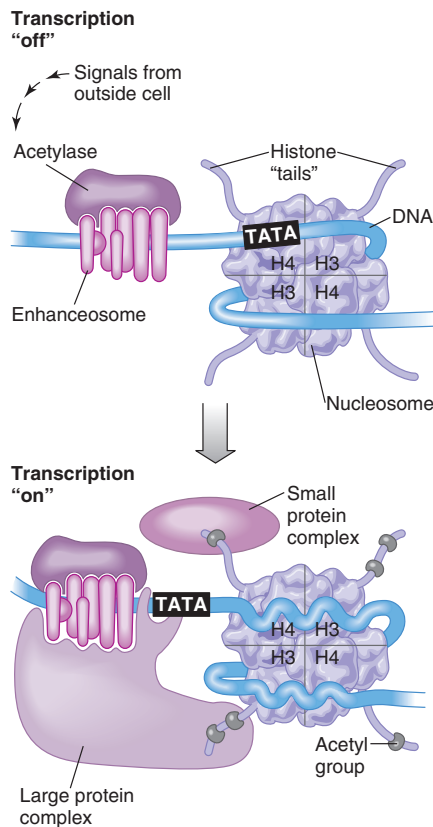


Figure 11.6 Acetylated histones allow transcription to begin. Once acetyl groups are added to particular amino acids in the tails of certain histones, the TATA box becomes accessible to transcription factors. H3 and H4 are histone types.

MicroRNAs

Chromatin remodeling determines which genes are transcribed. MicroRNAs act later in gene expression, preventing the translation of mRNA transcripts into protein. If chromatin remodeling is considered as an on/off switch to transcription,

then the control of microRNAs is more like that of a dimmer switch—fine-tuning gene expression at a later stage. **Figure 11.7** schematically compares chromatin modeling and the actions of microRNAs, and **figure 11.8** places them in the overall flow of genetic information.

MicroRNAs belong to a class of RNA molecules called noncoding RNAs, so-named because they were not among the first three major classes of RNA described (mRNA, tRNA, and rRNA). MicroRNAs are so small—just 21 or 22 bases long—that for many years, researchers unwittingly threw them out when doing experiments searching for longer molecules. In the cell, microRNAs are cut from precursors.

The human genome has close to 1,000 distinct microRNAs that regulate at least one-third of the protein-encoding genes. The DNA sequences that encode microRNAs are found in protein-encoding parts of the genome as well as in the vast regions that do not encode protein and are less well understood. From 1,000 to 200,000 microRNAs are present in a typical human cell.

Each type of microRNA binds to parts of the initial control regions (corresponding to DNA promoters) of a particular set of mRNAs, by complementary base pairing. When a microRNA binds a “target” mRNA, it prevents translation. Because a single type of microRNA has many targets, it effectively controls the expression of sets of genes at a time. In turn, a single type of mRNA can bind several different microRNAs. Researchers use computational tools (bioinformatics) to track, compare, and analyze all of these complex interactions.

Within the patterns of microRNA function may lie clues to developing new ways to fight disease, because these controls of gene expression have stood the test of evolutionary time. That is, the mRNAs that a microRNA targets may represent genes that function together. When one gene is abnormal, it impacts others, perhaps accounting for some cases of genetic heterogeneity (mutations in different genes causing the same phenotype).

The first practical applications of microRNAs are in cancer, because certain microRNAs are either more or less abundant in cancer cells than in healthy cells of the same type from which the cancer cells formed. Restoring the levels of microRNAs that normally suppress the too-rapid cell cycling of cancer, or blocking production of microRNAs that are too abundant in cancer,

Table 11.1 Disorders of Chromatin Remodeling				
Disease	MIM	Protein	Symptoms	Defect
α -thalassemia mental retardation syndrome	301040	ATRX	Anemia, mental retardation	Undermethylation of heterochromatin
ICF syndrome	242860	DNMT3B	Immunodeficiency, unstable centromeres, facial anomalies	Undermethylation of DNA repeats
Rett syndrome	312750	MECP2	Repetitive movements, irregular breathing, seizures, loss of motor control, neurodegeneration	Failure to remove acetyls from histones on gene <i>DLX5</i> expressed in brain
Rubinstein-Taybi syndrome	180849	CBP	Mental retardation, short stature, facial anomalies	Adds acetyl groups to certain histones, causing inappropriate transcription of some genes

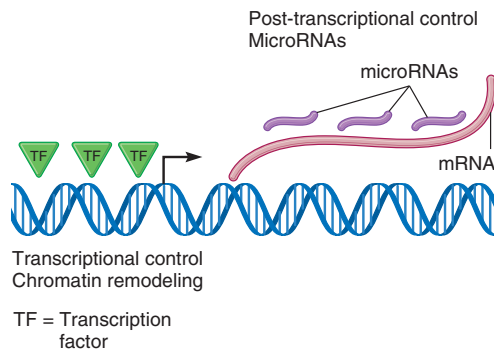


Figure 11.7 Control of gene expression. Chromatin opens to allow transcription factors to bind, whereas microRNAs bind to specific mRNAs, blocking their translation into protein.

could help to return cells to normal. In a related technology called RNA interference (RNAi), small, synthetic, double-stranded RNA molecules are introduced into selected cell types. Here they block gene expression in the same manner as the naturally-occurring microRNAs. Chapter 19 discusses RNAi further.

Key Concepts

1. Acetyl, phosphate, and methyl groups bind to histone proteins, controlling transcription.
2. Acetyl and phosphate groups turn on transcription; methyl groups turn it off.
3. The patterns of histones that activate a particular gene may follow rules like a code.
4. MicroRNAs bind to the control regions of specific mRNAs, blocking translation.

11.3 Maximizing Genetic Information

One of the marvels of the human genome is that it maximizes information—20,325 or so genes encode about 100,000 mRNAs, which in turn specify more than a million proteins. Figure 11.8 depicts this increase in information from gene to RNA to protein on the left, and the mechanisms that maximize the information on the right.

Several events account for the fact that proteins outnumber genes. The “genes in pieces” pattern of exons and introns and alternate splicing make this possible, much as a few items of clothing can be assembled into many outfits by combining them in different ways (**figure 11.9**). The different proteins that result from different uses of the information in a gene are called isoforms. That is, the exons of a gene can encode more than one protein. Which one or ones are transcribed depends upon circumstances. For example, when an infection begins, an immune system cell first secretes a short version of an antibody molecule that is presented on the cell’s surface. This alerts other cells. As the infection progresses, the cell transcribes an additional exon that extends the antibody in a way that enables it to be secreted into the bloodstream, where it attacks the pathogen.

Alternate splicing explains how a long sequence of DNA can specify more mRNAs than genes. On a part of chromosome 22, for example, 245 genes yield 642 mRNA transcripts. About 90 percent of all human genes are alternately spliced.

A DNA sequence that is an intron in one context may encode protein in another. Consider prostate specific antigen (PSA). This is a protein on certain cell surfaces that is overproduced in some prostate cancers. The gene for PSA has five exons and four introns. However, it is alternately

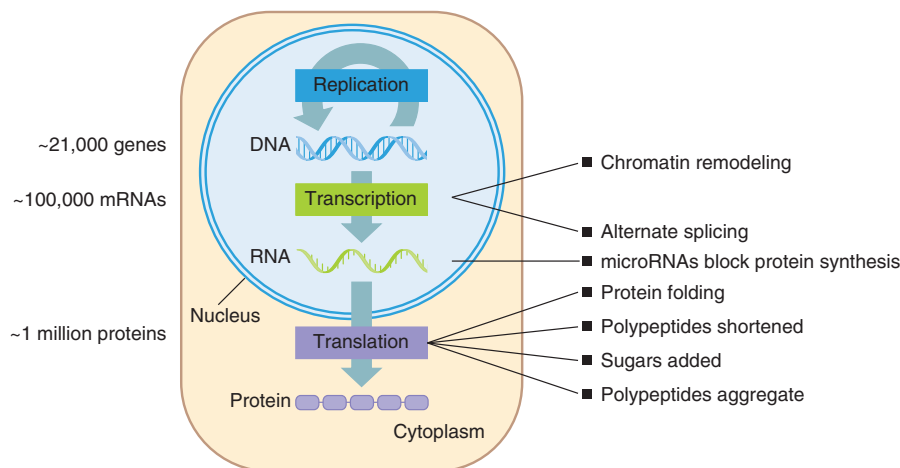


Figure 11.8 A summary of the events of gene expression. At the level of transcription, chromatin remodeling determines which genes are transcribed. Alternate splicing creates different forms of a protein by combining exons in different ways. MicroRNAs bind to the 3’ ends of mRNAs by complementary base pairing, blocking translation. At the level of translation, a protein must fold a certain way. Certain polypeptides must be shortened, attached to sugars, or aggregated.

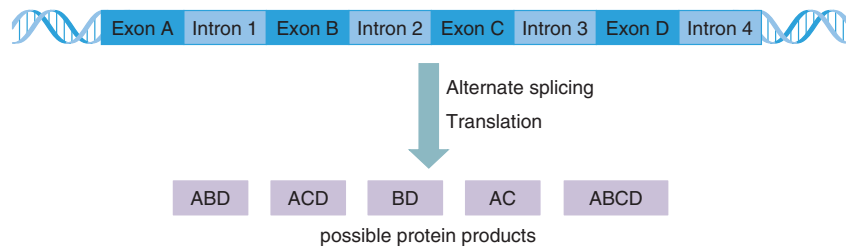


Figure 11.9 Exons provide flexibility in gene structure that expands gene number. Alternate splicing enables a cell to manufacture different versions of a protein by adding or deleting parts. Introns are removed and exons retained.

spliced to encode a different protein, called PSA-linked molecule (PSA-LM), that consists of the first exon and the fourth intron. The two proteins (PSA and PSA-LM) seem to work against each other. When the level of one is high, the other is low. New blood tests may consider levels of both proteins to more accurately assess the risk of developing prostate cancer.

Another way that introns may increase the number of proteins compared to genes is that a DNA sequence that is an intron in one gene's template strand may encode protein on the coding strand. This is the case for the gene for neurofibromin, which when mutant causes neurofibromatosis type 1, an autosomal dominant condition that causes benign tumors beneath the skin and spots on the skin surface. Within an intron of the neurofibromin gene, but on the coding strand, are instructions for three other genes.

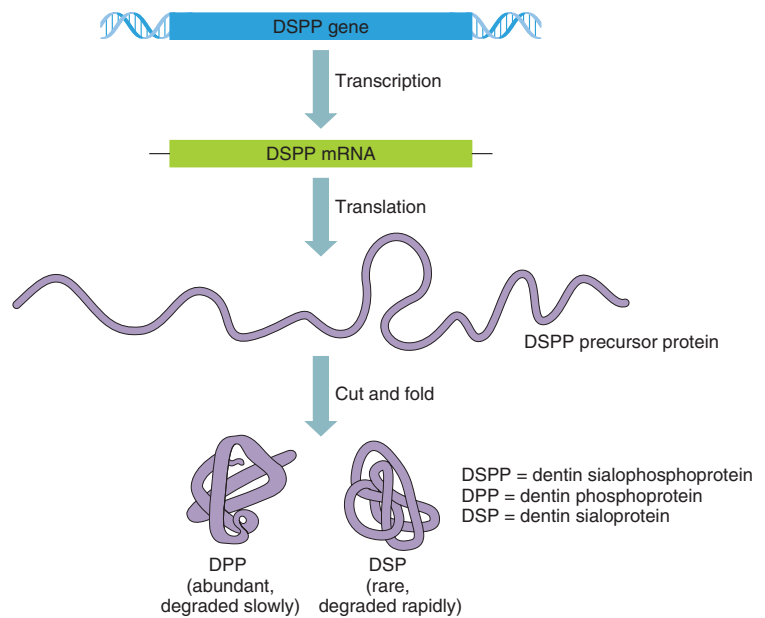
Reading a DNA sequence in two directions may be more commonplace than was thought—it might not have occurred to researchers to seek meaning in the complementary DNA strand of a sequence found to encode a protein. Finding dual meaning is a little like reading a novel backwards and discovering a second story! Such “bidirectional transcription” is seen in a condition called Huntington disease-like 2 (*HDL2*; MIM 606438). This disease causes the same symptoms as those seen in Huntington disease (HD), discussed in chapter 4. Also like HD, it is the result of a mutation that adds extra DNA triplets to a gene. The complement of part of the repeated region in *HDL2* is an exon of a gene called *junctophilin-3* (MIM 605268), which encodes a protein that anchors the endoplasmic reticulum to the plasma membrane.

At the level of translation, information is maximized when a protein is modified into different forms by the addition of sugars or lipids to create glycoproteins and glycolipids. Another way that one gene can encode more than one protein is if the protein is cut, yielding two products. This happens in dentinogenesis imperfecta (MIM 125490), which is an autosomal dominant condition that causes discolored, misshapen teeth with peeling enamel (figure 11.10). The dentin, which is the bonelike substance beneath the

enamel that forms the bulk of the tooth, is abnormal. Dentin is a complex mixture of extracellular matrix proteins, 90 percent of which are collagens. However, two proteins are unique to



a.



b.

Figure 11.10 Another way to encode two genes in one. (a) The misshapen, discolored, and enamel-stripped teeth of a person with dentinogenesis imperfecta were at first associated with deficiency of the protein DPP. Then researchers discovered that DSP is deficient, too, but is very scant. (b) Both DPP and DSP are cut from the same larger protein, but DSP is degraded faster.

dentin: the abundant dentin phosphoprotein (DPP) and the rare dentin sialoprotein (DSP). A single gene encodes both. DPP and DSP are translated from a single mRNA molecule as the precursor protein dentin sialophosphoprotein (DSPP). DPP is more abundant because DSP is degraded much faster.

Key Concepts

1. Only a tiny proportion of the genome encodes protein, yet the number of proteins greatly outnumbers known protein-encoding genes.
2. Alternate splicing, introns that encode protein, and cutting a precursor protein maximize the number of proteins that DNA encodes.

11.4 Most of the Human Genome Does Not Encode Protein

When the first molecular geneticists worked out the details of transcription and translation in the 1960s, they never imagined that only 1.5 percent of human DNA encodes protein. What does the “other” 98.5 percent do? It includes viral sequences, sequences that encode RNAs other than mRNA (called noncoding or ncRNAs), introns, promoters and other control sequences, and repeated sequences (**table 11.2**). In fact, most of the genome is transcribed—it isn’t “junk.” However, we do not yet know the functions of many transcripts.

Viral DNA

Our genomes include DNA sequences that represent viruses. Viruses are nonliving infectious particles that consist of a nucleic acid (DNA or RNA) encased in a coat built of proteins (see Reading 17.1). A virus replicates using a cell’s transcriptional and translational machinery to mass-produce viruses, which exit the cell. Sometimes the viral nucleic acid remains in a chromosome of a host cell. A DNA virus may do so directly; an RNA virus first uses an enzyme (reverse transcriptase) to copy its genetic material into DNA, which then inserts into a host chromosome.

About 100,000 sequences in our DNA, of varying length and comprising about 8 percent of the genome, were once a type of RNA virus called a retrovirus. The name refers to a retrovirus’ direction of genetic information transfer, which is opposite DNA to RNA to protein. Retroviral sequences in our chromosomes are termed “endogenous” because they are carried from generation to generation of the host, rather than acquired as an acute infection. The retroviruses whose genetic material is in our chromosomes are called human endogenous retroviruses, or HERVs.

By comparing HERV sequences to similar viruses in other primates, researchers have traced HERVs to a sequence

representing a virus that infected our ancestors’ genomes about 5 million years ago. Since then, HERV sequences have exchanged parts (recombined) and mutated to the extent that they no longer make us sick—yet they retain the theoretical ability to do so by recombining and mutating. In fact, researchers have combined parts of HERVs and regenerated an ancient, infectious HERV. They named it Phoenix, after the legendary bird that arose from the ashes. Harmless HERVs silently pass from human generation to generation because they are parts of our chromosomes. They tend to increase in number with time, as **figure 11.11** shows.

Noncoding RNAs

When researchers began to annotate, or describe the function, of regions of the human genome, they were surprised to find that much more of the genome is transcribed than would be predicted based on the number and diversity of proteins that a human body can produce. The emerging explanation is that the human genome “instruction manual” is much more extensive than anyone realized. In fact, nearly all of the human genome can be transcribed, and much of it is in the form of noncoding RNAs (ncRNAs), which include all types except the mRNAs. The best-studied are the tRNAs and rRNAs, simply because they were discovered first, as the genetic code was cracked and protein synthesis described in the 1960s. More recently, several other classes of RNAs have been discovered, and they are summarized in **table 11.2**.

The rate of transcription of a cell’s tRNA genes is attuned to cell specialization. The proteins of a skeletal muscle cell, for example, would require different amounts of certain amino acids than the proteins of a white blood cell, and therefore different amounts of the corresponding tRNAs, too. Human tRNA genes are dispersed among the chromosomes in clusters—25 percent of them are on the sixth-largest chromosome, for example. Altogether, our 500 or so types of tRNA genes account for 0.1 percent of the genome.

The 243 types of rRNA genes are grouped on six chromosomes, each cluster harboring 150 to 200 copies of a 44,000-base repeat sequence. Once transcribed from these clustered genes, the rRNAs go to the nucleolus, where another type of ncRNA called small nucleolar RNA (snoRNA) cuts them into their final forms.

Hundreds of thousands of ncRNAs are neither tRNA nor rRNA, nor snoRNAs, nor microRNAs, nor the other less abundant types described in **table 11.2**. Instead, they are transcribed from DNA sequences called **pseudogenes**. A pseudogene is very similar in sequence to a particular protein-encoding gene, and it may be transcribed into RNA, but it is not translated into protein. Presumably a pseudogene is altered in sequence from an ancestral gene in a way that impairs its translation—perhaps the encoded amino acids cannot fold into a functional protein. Pseudogenes may be remnants of genes past, variants that diverged from the normal sequence too greatly to encode a working protein. Pseudogenes are incredibly common in the human genome.

Table 11.2 Some Nonprotein-Encoding Parts of the Human Genome

Type of Sequence	Function or Characteristic
Viral DNA	Evidence of past infection
Noncoding RNA genes	
tRNA genes	Connect mRNA codon to amino acid
rRNA genes	Parts of ribosomes
Pseudogenes	DNA sequences very similar to known genes that are not translated
Small nucleolar RNAs (snoRNA)	Process rRNA in nucleolus
Small nuclear RNAs (snRNA)	Parts of spliceosomes
Telomerase RNA	Adds bases to chromosome tips
Xist RNA	Inactivates one X chromosome in cells of females
Introns	Parts of genes that are cut out of mRNA
Promoters and other control sequences	Guide enzymes that carry out DNA replication, transcription, or translation
Small interfering RNAs (siRNA)	Control transcription
MicroRNAs (miRNA)	Control transcription of many genes
Repeats	
Transposons	Repeats that move around the genome
Telomeres	Chromosome tips that control the cell cycle
Centromeres	Largest constriction in a chromosome, providing attachment points for spindle fibers
Duplications of 10 to 300 kilobases	Unknown
Simple short repeats	Unknown

amino acid sequence. Perhaps repeat size or number constitute another type of molecular language. Or, perhaps some types of repeats help to hold a chromosome together.

The most abundant type of repeat is a sequence of DNA that can jump about the genome. It is called a transposable element, or **transposon** for short. Geneticist Barbara McClintock originally identified transposons in corn in the 1940s, and they were rediscovered in bacteria in the 1960s. Transposons comprise about 45 percent of the human genome sequence, and typically repeat in many copies. Some transposons include parts that encode enzymes that cut them out of one chromosomal site and integrate them into another. Transposons are classified into four groups: by size, whether they are transcribed into RNA, which enzymes they use to move, and whether they resemble bacterial transposons.

An example of a specific type of repeat is an Alu sequence. Each Alu repeat is about 300 bases long, and a human genome may contain 300,000 to 500,000 of them. Alu repeats comprise 2 to 3 percent of the genome, and they have been increasing in number over time because they can copy themselves. We don't know exactly what these common repeats do, if anything. They may serve as attachment points for proteins that bind newly replicated DNA to parental strands before anaphase of mitosis, when replicated chromosomes pull apart.

Rarer classes of repeats include those that comprise telomeres, centromeres, and rRNA gene clusters; duplications of 10,000 to 300,000 bases (10 to 300 kilobases); cop-

ies of pseudogenes; and simple repeats of one, two, or three bases. In fact, the entire human genome may have duplicated once or even twice, as is discussed further in chapter 16.

Repeats

The human genome is riddled with highly repetitive sequences that may be a different type of information than a protein's

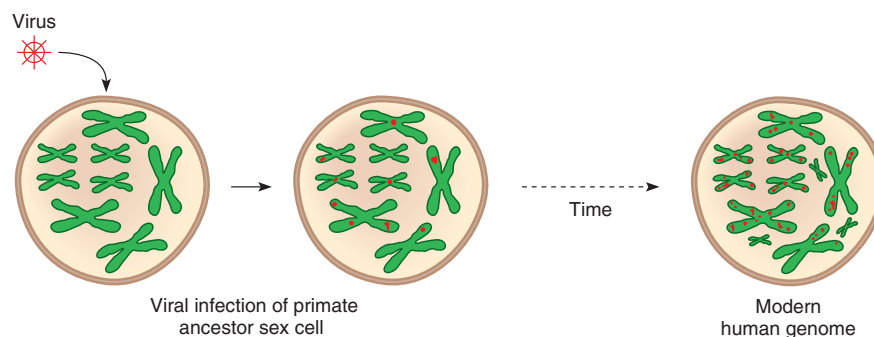


Figure 11.11 The human genome includes viral DNA sequences. Most if not all of them do not harm us.

Our understanding of the functions of repeats lags far behind our knowledge of the roles of the various noncoding RNA genes. Repeats may make sense in light of evolution, past and future. Pseudogenes are likely vestiges of genes that functioned in our nonhuman ancestors. Perhaps the repeats that seem to have no obvious function today will serve as raw material from which future genes may arise.

Discovery of the intricate controls of gene expression has led to a new definition of a gene, greatly expanded from the one-gene, one-protein idea of years past. A gene is a DNA sequence that contributes to a phenotype or function, plus the sequences,

both in the gene and outside it, that control its expression. The next chapter looks at the different types of changes in the DNA sequence—mutations—and their consequences.

Key Concepts

1. Most of the genome encodes many types of RNA as well as viral sequences, introns, promoters, and other control sequences and repeats.
2. We do not know the functions of some repeats.

Summary

11.1 Gene Expression Through Time and Tissue

1. Changes in gene expression occur over time at the molecular level (globin switching), at the tissue level (blood plasma), and at the organ/gland level (pancreas development).
2. **Proteomics** catalogs the types of proteins in particular cells, tissues, organs, or entire organisms under specified conditions.

11.2 Control of Gene Expression

3. Acetylation of certain histones enables the transcription of associated genes. Phosphorylation and methylation are also important in **chromatin remodeling**.
4. MicroRNAs bind to certain mRNAs, blocking translation.

11.3 Maximizing Genetic Information

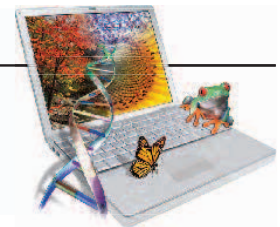
5. A small part of the genome encodes protein, but these genes specify a much greater number of proteins.
6. Alternate splicing, use of introns, and cutting proteins translated from a single gene contribute to protein diversity.

11.4 Most of the Human Genome Does Not Encode Protein

7. The nonprotein-encoding part of the genome includes viral sequences, noncoding RNAs, introns, promoters and other controls, and repeats.

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Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

1. Why is control of gene expression necessary?
2. Define *epigenetics*.
3. Distinguish between the type of information that epigenetics provides and the information in the DNA sequence of a protein-encoding gene.
4. Describe three types of cells and how they differ in gene expression from each other.
5. Explain how a mutation in a promoter can affect gene expression.
6. What is the environmental signal that stimulates globin switching?
7. How does development of the pancreas illustrate differential gene expression?
8. How do histones control gene expression, yet genes also control histones?
9. Name a mechanism that silences transcription of a gene and a mechanism that blocks translation of an mRNA.
10. What controls whether histones enable DNA wrapped around them to be transcribed?
11. What are two ways that microRNA functioning is complex?
12. Describe three ways that the number of proteins exceeds the number of protein-encoding genes in the human genome.
13. How can alternate splicing generate more than one type of protein from the information in a gene?
14. In the 1960s, a gene was defined as a continuous sequence of DNA, located permanently at one place on a chromosome, that specifies a sequence of amino acids from one strand. List three ways this definition has changed.
15. Give an example of a discovery mentioned in the chapter that changed the way we think about the genome.

Applied Questions

- Several new drugs inhibit the enzymes that either put acetyl groups on histones or take them off. Would a drug that combats a cancer caused by too little expression of a gene that normally suppresses cell division add or remove acetyl groups?
- Chromosome 7 has 863 protein-encoding genes, but many more proteins. The average gene is 69,877 bases, but the average mRNA is 2,639 bases. Explain both of these observations.
- CHARGE syndrome (MIM 214800) causes heart defects, visual problems, facial palsy, blocked nostrils, and difficulty swallowing. A mutation in a gene called *Chd1* causes the condition. The protein product of this gene recognizes and binds methyl groups on certain histones. Explain how this mutation leads to pleiotropy (multiple symptoms).
- How many different proteins encompassing two exons can be produced from a gene that has three exons?
- Many people with trisomy 21 Down syndrome (an extra chromosome 21; see section 13.3) who survive into adulthood develop early-onset Alzheimer disease. The APP gene which when mutant causes this form of Alzheimer disease is on chromosome 21. Explain how this form of Alzheimer disease in trisomy 21 individuals differs from the same disorder caused by a mutation in the APP promoter in a person who has the normal two copies of chromosome 21.

Web Activities

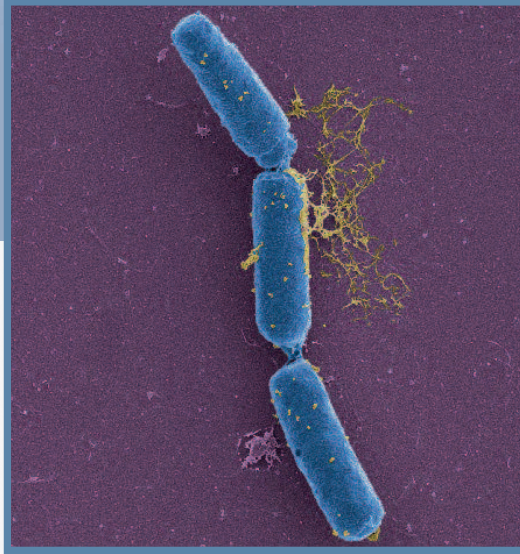
- Gene expression profiling tests began to be marketed just a few years ago. Google "Oncotype DX," "MammaPrint," or simply "gene expression profiling in cancer" and describe how classifying a particular cancer based on gene expression profiling can improve diagnosis and/or treatment. (Or apply this question to a different type of disease.)

Case Studies and Research Results

- Jerrold is 38 years old. His body produces too much of the hormone estrogen, which has enlarged his breasts. He had a growth spurt and developed pubic hair by age 5, and then his growth dramatically slowed so that his adult height is well below normal. He has a very high-pitched voice and no facial hair, which reflect the excess estrogen. Jerrold's son, Timmy, is 8 years old and has the same symptoms.
Jerrold and Timmy have an overactive gene for aromatase, an enzyme required to synthesize estrogen. Five promoters control expression of the gene in different tissues, and each promoter is activated by a different combination of hormonal signals. The five promoters lead to estrogen production in skin, fat, brain, gonads (ovaries and testes) and placenta. In premenopausal women, the ovary-specific promoter is highly active, and estrogen is abundant. In men and postmenopausal women, however, only small amounts of estrogen are normally produced, in skin and fat. The father and son have a wild type aromatase gene, but high levels of estrogen in several tissues, particularly fat, skin, and blood. They do, however, have a mutation that turns around an adjacent gene so that the aromatase gene falls under the control of a different promoter. Suggest how this phenotype arises.
- Margaret is 102 years old, and she still walks at least half a mile a day, albeit slowly. She is a trim vegetarian who has rarely been ill her entire life. Morris is an obese, balding 62-year-old man who has high blood pressure and colon cancer. How might their proteome portraits, such as the one in figure 11.4, differ? (Hint: Reread Reading 3.1, Genes and Longevity.)

Forensics Focus

- Establishing time of death is critical information in a murder investigation. Forensic entomologists can estimate the "postmortem interval" (PMI), or the time at which insects began to deposit eggs on the corpse, by sampling larvae of specific insect species and consulting developmental charts to determine the stage. The investigators then count the hours backwards to estimate the PMI. Blow flies are often used for this purpose, but their three larval stages look remarkably alike in shape and color, and development rate varies with environmental conditions. With luck, researchers can count back 6 hours from the developmental time for the largest larvae to estimate the time of death.
In many cases, a window of 6 hours is not precise enough to narrow down suspects when the victim visited several places and interacted with many people in the hours before death. Suggest a way that gene expression profiling might be used to more precisely define the PMI and extrapolate a probable time of death.



Bacillus anthracis—anthrax—has been used as a bioweapon.

CHAPTER

12

Gene Mutation

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The Amerithrax Story

In the weeks following the 2001 terrorist attacks in the United States, several people, including two senators and individuals at several media outlets, received letters containing threatening messages and a mysterious white powder. Of the 22 people who opened the letters and were exposed to the contents—spores of *Bacillus anthracis*—five died of inhalation anthrax. The Federal Bureau of Investigation (FBI) dubbed the investigation, which took 7 years, “Amerithrax.”

Anthrax is a bacterial infection usually contracted through the skin from hoofed mammals (figure). Inhalation anthrax is extremely rare, and has been used as a bioweapon (see section 17.5). Within a year of the mysterious letters, scientists identified the Amerithrax anthrax as a particular laboratory strain called Ames, and had sequenced the genome of the bacterium that killed the first victim, a reporter at a tabloid newspaper. Old-fashioned sleuthing was required, too. Investigators collected and typed more than 1,000 strains of anthrax from all over the world, zeroing in on eight samples that matched the letter anthrax—they all came from a government lab in Maryland. There, an experienced technician studied the suspect bacteria and noted that they formed colonies that looked unusual—they were mutants.

The strain of anthrax that had been sent to the victims bore five distinctive genetic changes, or mutations, that served as an identity tag. It was named “RMR-1029.” Over a 2-year period, FBI investigators found the only person who had access to these bacteria—a researcher who reportedly committed suicide in August 2008, just before he was to be indicted. The case ended with his death, but questions remain.

During the Amerithrax scare, the author of this book also received a seemingly threatening letter, with peculiar block printing and strange messages. It came from Libya and was delivered to a doctor's office that sounded like the one where she is a genetic counselor. An assistant called the police and the FBI came and picked up the letter, keeping it for months, as all who touched it took antibiotics. The letter was finally returned, triple-bagged with poison symbols, deemed safe. The message within was from a genetics instructor politely correcting an error in this book!

12.1 The Nature of Mutations

A **mutation**, used as a noun, is a change in a DNA sequence that is present in less than 1 percent of individuals in a population. Used as a verb, “mutation” or “mutate” refers to the process of altering a DNA sequence. Mutations range in size from alteration of a single DNA base; to deletion or duplication of tens, hundreds, thousands, or even millions of bases; to missing or extra entire chromosomes. This chapter discusses smaller-scale mutations, and chapter 13 considers mutation at the chromosomal level.

Mutation can affect any part of the genome, including sequences that encode proteins or control transcription; in introns; repeats; and sites critical to intron removal and exon splicing. However, not all DNA sequences are equally likely to mutate.

The effects of mutation vary. Mutations may impair a function, have no effect, or even be beneficial. A deleterious (harmful) mutation can stop or slow production of a protein, overproduce it, or impair the protein's function—such as altering its secretion, location, or interaction with another protein. Geneticists describe the effect of a mutation as a “loss of function” when the gene's product is reduced or absent, or as a “gain of function” when the gene's action changes in some way. Most mutations are recessive and cause a loss of function. Gain-of-function mutations tend to be dominant.

The terms *mutation* and *polymorphism* can be confusing, because both denote a genetic change. Recall from chapter 7 that a single nucleotide polymorphism, or SNP, is a single base change. So are many mutations. The distinction between mutation and polymorphism is largely artificial. Geneticists define a mutation as being present in less than 1 percent of a population and a polymorphism as being present in more than 1 percent of a population. The consequences of this distinction provide the logic underlying the more popular view that a mutation is “bad” but a polymorphism is a “harmless variant.” That is, if a genetic change greatly impairs health, individuals with it are unlikely to reproduce, and the mutant allele remains uncommon. (Chapter 15 explains why it doesn't actually disappear.) A polymorphism that does not harm health, elevates risk of illness only slightly, or is even beneficial, will remain prevalent in a population or even increase in frequency. This chapter deals mostly with mutations that alter the phenotype.

Not all mutations are harmful, in contrast to their depiction in science fiction. For example, a mutation protects against HIV infection. About 1 percent of the general population is homozygous for a recessive allele that encodes a cell surface protein called CCR5 (see figure 17.12). To infect an immune system cell, HIV must bind CCR5 and another protein. Because the mutation prevents CCR5 from moving to the cell surface, HIV cannot bind. Heterozygotes are partially protected against HIV infection.

The term *mutation* refers to genotype—that is, a change at the DNA or chromosome level. The familiar term **mutant** refers to phenotype. The nature of a mutant phenotype depends upon how the mutation affects the gene's product or activity, and usually connotes an abnormal or unusual characteristic. However, a mutant phenotype may also be an uncommon variant that is nevertheless “normal,” such as red hair.

In an evolutionary sense, mutation has been essential to life, because it produces individuals with variant phenotypes who are better able to survive specific environmental challenges, including illnesses. Our evolutionary relatedness to other species enables us to study many mutations in nonhuman species, which can provide information on our own (**figure 12.1**).



a.



b.

Figure 12.1 Animal models of human diseases. (a) Mice that have been genetically modified to harbor human genes for neurological disorders, such as Huntington disease and Alzheimer disease, are tested on a rotating drum with a grooved surface called a Rotarod. As the animals lose motor skills, it becomes more difficult for them to hold onto an accelerating Rotarod. (b) This dog is a natural model of amyotrophic lateral sclerosis (Lou Gehrig's disease) in humans. Mutation in the same gene—superoxide dismutase 1—causes about 2 percent of human cases, as well as the canine cases.

A mutation may be present in all the cells of an individual or just in some cells. In a **germline mutation**, the change occurs during the DNA replication that precedes *meiosis*. The resulting gamete and all the cells that descend from it after fertilization have the mutation—that is, every cell in the body. In contrast, a **somatic mutation** happens during DNA replication before a *mitotic* cell division. All the cells that descend from the original changed cell are altered, but they might only comprise a small part of the body. Somatic mutations are more likely to occur in cells that divide often, such as hair root cells, because there are more opportunities for replication errors. Reading 18.1 and figure 18.5 concern somatic mutations and cancer.

12.2 Two Mutations

Identifying how a mutation causes symptoms has clinical applications, and also reveals the workings of biology. Following are two examples of mutations that cause disease.

The Beta Globin Gene Revisited

The first genetic illness understood at the molecular level was sickle cell disease (**figure 12.2**). In 1904, young medical intern Ernest Irons noted “many pear-shaped and elongated forms”

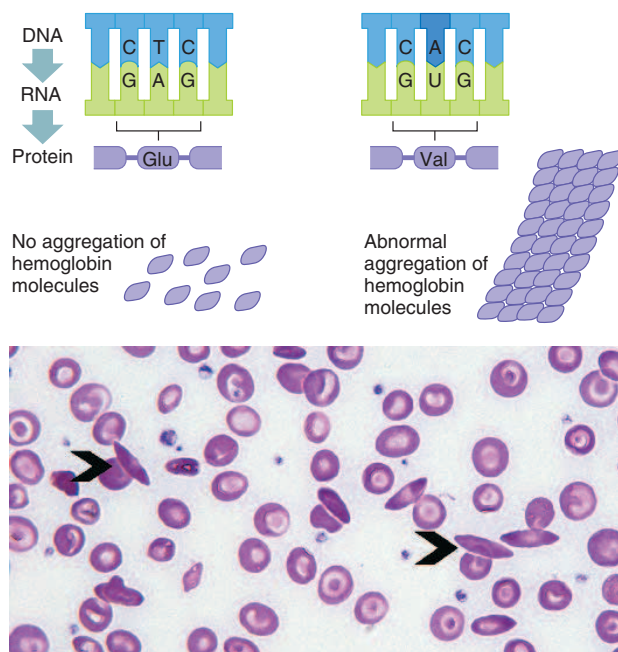


Figure 12.2 Sickle cell disease results from a single DNA base change that substitutes one amino acid in the protein (valine replaces glutamic acid). This changes the surfaces of the molecules, and they aggregate into long, curved rods that deform the red blood cell. Arrows in photograph point to sickled cells.

in a blood sample from a dental student in Chicago who had anemia. Irons sketched this first view of sickle cell disease at the cellular level, and reported his findings to his supervisor, physician James Herrick. Alas, Herrick published the work but did not mention Irons. Herrick has been credited with the discovery ever since.

In 1949, Linus Pauling discovered that hemoglobin from healthy people and from people with the anemia, when placed in a solution in an electrically charged field (a technique called electrophoresis), moved to different positions. Hemoglobin from the parents of people with the anemia, who were carriers, moved to both positions.

The difference between the two types of hemoglobin lay in beta globin. Recall from figure 11.1 that adult hemoglobin consists of two alpha polypeptide subunits and two beta subunits. Protein chemist V. M. Ingram took a shortcut to localize the mutation in the 146-amino-acid-long beta subunit. He cut normal and sickle hemoglobin with a protein-digesting enzyme, separated the pieces, stained them, and displayed them on filter paper. The patterns of fragments—known as peptide fingerprints—were different for the two types of hemoglobin. This meant, Ingram deduced, that the two molecules differ in amino acid sequence. Then he discovered the difference. One piece of the molecule in the fingerprint, fragment four, occupied a different position in each of the two types of hemoglobin. Because this peptide was only 8 amino acids long, Ingram needed to decipher only that short sequence to find the site of the mutation. It was a little like finding which sentence on a page contains a typographical error.

Ingram identified the tiny mutation responsible for sickle cell disease: a substitution of the amino acid valine for the glutamic acid that is normally the sixth amino acid in the beta globin polypeptide chain. At the DNA level, the change was even smaller—a CTC to a CAC, corresponding to RNA codons GAG and GUG. This was learned after researchers deciphered the genetic code. The valine at this position changes the surfaces of hemoglobin molecules so that in low-oxygen conditions they attach at many more points than they would if the wild type glutamic acid were at the site. The aggregated hemoglobin molecules form ropelike cables that bend red blood cells into rigid, fragile, sickle-shaped structures. The misshapen cells lodge in narrow blood vessels, cutting off local blood supplies. Once a blockage occurs, sickling speeds up and spreads, as the oxygen level falls. The result is great pain in the blocked body parts, particularly the hands, feet, and intestines. The bones ache, and depletion of normal red blood cells causes the great fatigue of anemia.

Sickle cell disease was the first inherited illness linked to a molecular abnormality, but it wasn't the first known condition that results from a mutation in the beta globin gene. In 1925, Thomas Cooley and Pearl Lee described severe anemia in Italian children, and in the decade following, others described a milder version of “Cooley’s anemia,” also in Italian children. The disease was named thalassemia, from the Greek for “sea,” in light of its high prevalence in the Mediterranean area. The two disorders turned out to be the same. The severe form, sometimes called thalassemia major, results from a homozygous

mutation in the beta globin gene. The milder form, called thalassemia minor, affects some individuals who are heterozygous for the mutation.

Once researchers had worked out the structure of hemoglobin, and learned that different globins function in the embryo and fetus, the molecular basis of thalassemia became clear. The disorder that is common in the Mediterranean is more accurately called beta thalassemia (MIM 141900), because the symptoms result from too few beta globin chains. Without them, not enough hemoglobin molecules are assembled to effectively deliver oxygen to tissues. Fatigue and bone pain arise during the first year of life as the child depletes fetal hemoglobin, and the “adult” beta globin genes are not transcribed and translated on schedule.

As severe beta thalassemia progresses, red blood cells die because the excess of alpha globin chains prevents formation of hemoglobin molecules. Liberated iron slowly destroys the heart, liver, and endocrine glands. Periodic blood transfusions can control the anemia, but they hasten iron buildup and organ damage. Drugs called chelators that entrap the iron can extend life past early adulthood, but they are very costly and not available in developing nations.

Disorders of Orderly Collagen

Much of the human body consists of the protein collagen, which is a major component of connective tissue. Collagen accounts for more than 60 percent of the protein in bone and cartilage and provides 50 to 90 percent of the dry weight of skin, ligaments, tendons, and the dentin of teeth. Collagen is in parts of the eyes and the blood vessel linings, and it separates cell types in tissues.

Genetic control of collagen synthesis and distribution is complex; more than thirty-five collagen genes encode more

than twenty types of collagen molecules. Other genes affect collagen, too. Mutations in the genes that encode collagen, not surprisingly, lead to a variety of medical problems (**table 12.1**). These disorders are particularly devastating, not only because collagen is nearly everywhere, but because collagen has an extremely precise conformation that is easily disrupted, even by slight alterations that might have little effect in proteins with other shapes (**figure 12.3**).

Collagen is sculpted from a longer precursor molecule called procollagen, which consists of many repeats of the amino acid sequence glycine-proline-modified proline. Three procollagen chains entwine. Two of the chains are identical and are encoded by one gene, and the other is encoded by a second gene and therefore has a different amino acid sequence. The electrical charges and interactions of these amino acids with water coil the procollagen chains into a very regular triple helix, with space in the middle only for tiny glycine. The ragged ends of the polypeptides are snipped off by enzymes to form mature collagen. The collagen fibrils continue to associate with each other outside the cell, building the fibrils and networks that hold the body together. So important is the precision of collagen formation that a mutation that controls placement of a single hydroxyl chemical group ($-OH^-$) on collagen causes a severe form of osteogenesis imperfecta (“brittle bone disease”), described in chapter 5 and shown in figure 3.21a.

The boy shown in **figure 12.4** has a form of Ehlers-Danlos syndrome. A mutation prevents his procollagen chains from being cut, and, as a result, collagen molecules cannot assemble. Instead, they form ribbonlike fibrils that lack the tensile strength to keep the skin from becoming too stretchy. Other collagen mutations cause missing procollagen chains, kinks in the triple helix, and defects in aggregation outside the cell.

Table 12.1 Some Collagen Disorders

Disorder	MIM	Genetic Defect (Genotype)	Signs and Symptoms (Phenotype)
Alport syndrome	203780	Mutation in type IV collagen interferes with tissue boundaries	Deafness and inflamed kidneys
Aortic aneurysm	100070	Missense mutation substitutes <i>arg</i> for <i>gly</i> in $\alpha 1$ gene	Aorta bursts
Chondrodysplasia	302950	Deletion, insertion, or missense mutation replaces <i>gly</i> with bulky amino acids	Stunted growth, deformed joints
Dystrophic epidermolysis bullosa	226600	Collagen fibrils that attach epidermis to dermis break down	Skin blisters on any touch
Ehlers-Danlos syndrome	130050	Missense mutations replace <i>gly</i> with bulky amino acids; deletions or missense mutations disrupt intron/exon splicing	Stretchy, easily scarred skin, lax joints
Osteoarthritis	165720	Missense mutation substitutes <i>cys</i> for <i>arg</i> in $\alpha 1$ gene	Painful joints
Osteogenesis imperfecta type I	166200	Inactivation of α allele reduces collagen triple helices by 50%	Easily broken bones; blue eye whites; deafness
Stickler syndrome	108300	Nonsense mutation in procollagen	Joint pain, degeneration of vitreous gel and retina

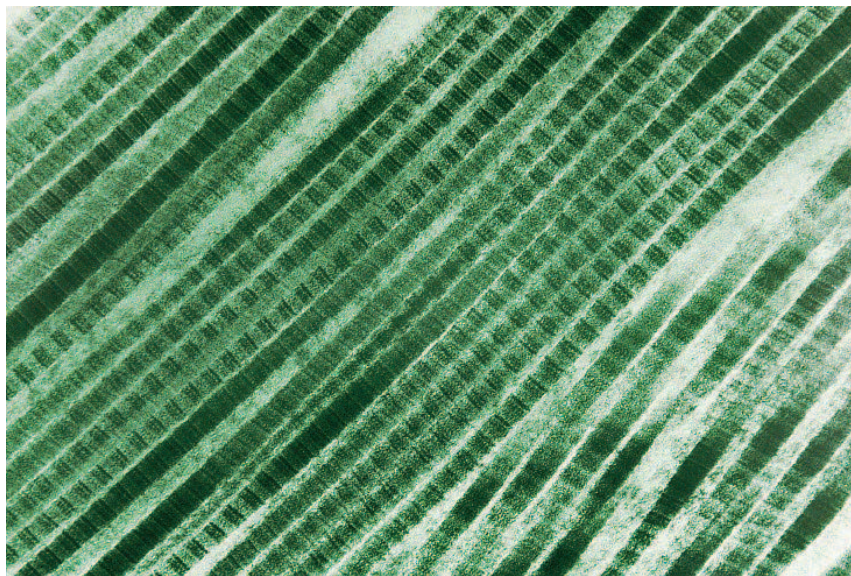
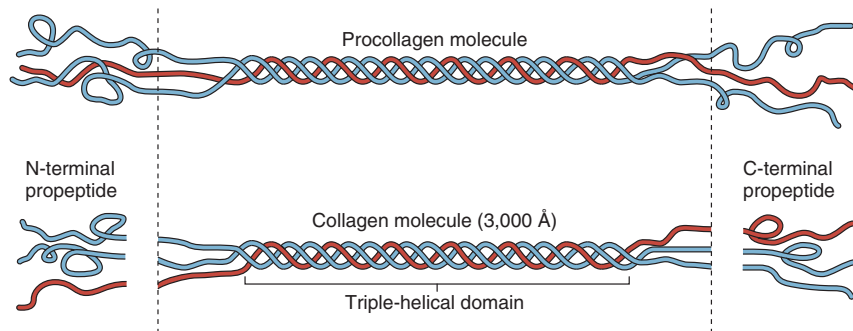


Figure 12.3 Collagen has a very precise conformation. The α_1 collagen gene encodes the two blue polypeptide chains, and the α_2 procollagen gene encodes the third (red) chain. The procollagen triple helix is shortened before it becomes functional, forming the fibrils and networks that comprise much of the human body.



Figure 12.4 A disorder of connective tissue produces stretchy skin. A mutation that blocks trimming of procollagen chains to produce collagen causes the stretchy skin of Ehlers-Danlos syndrome type I.

Aortic aneurysm is a serious connective tissue disorder. It is part of Marfan syndrome. Detection of mutations that cause Marfan syndrome before symptoms arise can be lifesaving, because frequent ultrasound exams can detect aortic weakening early enough to patch the vessel before it bursts. **Table 12.2** describes how a few other mutations impair health.

One Disorder or Several?

Geneticists can be inconsistent when assigning disease names to mutations. For one gene, different mutations may cause differing degrees or different subsets of symptoms of one syndrome. Yet for another gene, different mutations cause different disorders. For example, all mutations in the *CFTR* gene cause cystic fibrosis—whether that includes the full spectrum of impaired breathing and digestion, or just male infertility. CF can affect different tissues in different individuals. Yet different mutations in the beta globin gene cause the clinically distinct sickle cell disease and beta thalassemia, two different disorders, although

Table 12.2 How Mutations Cause Disease

Disorder	MIM	Protein	Genetic Defect (Genotype)	Signs and Symptoms (Phenotype)
Cystic fibrosis	602421	Cystic fibrosis transmembrane regulator (CFTR)	Missing amino acid or other defect alters conformation of chloride channels in certain epithelial cell plasma membranes. Water enters cells, drying out secretions.	Frequent lung infection, pancreatic insufficiency
Duchenne muscular dystrophy	310200	Dystrophin	Deletion eliminates dystrophin, which normally binds to inner face of muscle cell plasma membranes, maintaining cellular integrity. Cells and muscles weaken.	Gradual loss of muscle function
Familial hypercholesterolemia	143890	LDL receptor	Deficient LDL receptors cause cholesterol to accumulate in blood.	High blood cholesterol, early heart disease
Hemophilia A	306700	Factor VIII	Absent or deficient clotting factor causes hard-to-control bleeding.	Slow or absent blood clotting
Huntington disease	143100	Huntingtin	Extra bases in the gene add amino acids to the protein product, which impairs certain transcription factors and proteasomes.	Uncontrollable movements, personality changes
Marfan syndrome	154700	Fibrillin or transforming growth factor β receptor	Deficient connective tissue protein in lens and aorta.	Long limbs, weakened aorta, spindly fingers, sunken chest, lens dislocation
Neurofibromatosis type 1	162200	Neurofibromin	Defect in protein that normally suppresses activity of a gene that causes cell division.	Benign tumors of nervous tissue beneath skin

they both affect part of the same tissue, blood, and are therefore lumped together as “hemoglobinopathies.”

Adding to the inconsistency is a gene such as *lamin A*. Unlike *CFTR*, in which mutations cause variations of one disease, and beta globin, in which mutations cause different diseases of the same tissue, mutations in *lamin A* cause different disorders that affect very different tissues. *Lamin A* mutations cause the rapid-aging disorder Hutchinson-Gilford progeria syndrome (see figure 3.22 and table 3.4), and at least six other conditions, including muscular dystrophies and a heart disorder. Lamin A proteins form a network beneath the inner nuclear membrane that interacts with chromatin. Apparently different mutations affect *lamin A*'s interactions with chromatin in ways that cause the diverse associated disorders.

Key Concepts

1. Mutations add, delete, or rearrange genetic material in a germline cell or somatic cell.
2. In sickle cell disease, a mutation causes hemoglobin to crystallize in a low-oxygen environment, bending red blood cells into sickle shapes and impairing circulation. In beta thalassemia, beta globin is absent or scarce, depleting hemoglobin molecules.
3. Mutations in collagen genes can disrupt the protein's precise organization.
4. Mutations in a gene may cause either different versions of the same disease or distinct illnesses.

12.3 Causes of Mutation

A mutation can occur spontaneously or be induced by exposure to a chemical or radiation. An agent that causes mutation is called a **mutagen**.

Spontaneous Mutation

A spontaneous mutation can be a surprise. For example, two healthy people of normal height may have a child with achondroplasia, an autosomal dominant form of dwarfism. How could this happen when no other family members are affected? If the mutation is dominant, why are the parents of normal height? The child has a genetic condition, but he did not inherit it. Instead, he originated it. His siblings have no higher risk of inheriting the condition than anyone in the general population, but each of his children will face a 50 percent chance of inheriting it. The boy's achondroplasia arose from a *de novo*, or new, mutation in a parent's gamete. This is a spontaneous mutation—that is, it is not caused by a mutagen. A spontaneous mutation usually originates as an error in DNA replication.

One cause of spontaneous mutation stems from the chemical tendency of free nitrogenous bases to exist in two slightly different structures, called tautomers. For extremely short times, each base is in an unstable tautomeric form. If, by chance, such an unstable base is inserted into newly forming DNA, an error will be generated and perpetuated when that strand replicates. **Figure 12.5** shows how this can happen.

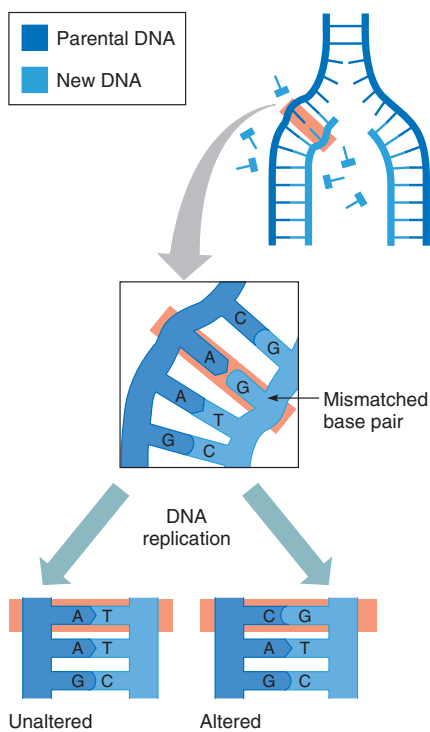


Figure 12.5 Spontaneous mutation. DNA bases are very slightly chemically unstable, and fleetingly they exist in alternate forms. If a replication fork encounters a base in its unstable form, a mismatched base pair can result. After another round of replication, one of the daughter cells has a different base pair than the one in the corresponding position in the original DNA. (This figure depicts two rounds of DNA replication.)

Spontaneous Mutation Rate

The spontaneous mutation rate varies for different genes. The gene that, when mutant, causes neurofibromatosis type 1 (NF1), for example, has a very high mutation rate, arising in 40 to 100 of every million gametes (table 12.3). NF1 affects 1 in 3,000 births, about half in families with no prior cases. The gene's large size may contribute to its high mutability—there are more ways for its sequence to change, just as there are more opportunities for a misspelling to occur in a long sentence than in a short one.

Each human gene has about a 1 in 100,000 chance of mutating and each of us probably carries a few new spontaneously mutated genes. Mitochondrial genes mutate at a higher rate than nuclear genes because they cannot repair DNA (see section 12.6.)

Estimates of the spontaneous mutation rate for a particular gene are usually derived from observations of new, dominant conditions, such as achondroplasia in the boy. This is possible because a new dominant mutation is detectable simply by observing the phenotype. In contrast, a new recessive mutation would not be obvious until two heterozygotes produced a homozygous recessive offspring with a noticeable phenotype.

The spontaneous mutation rate for autosomal genes can be estimated using the formula: number of *de novo* cases/2X, where X is the number of individuals examined. The denominator has a factor of 2 to account for the nonmutated homologous chromosome.

Spontaneous mutation rates in human genes are difficult to assess because our generation time is long—usually 20 to 30 years. In bacteria, a new generation arises every half hour or so, and mutation is therefore much more frequent. The genetic material of viruses also spontaneously mutates rapidly.

Table 12.3 Mutation Rates of Some Genes That Cause Inherited Disease

Disorder	MIM	Mutations per Million Gametes	Signs and Symptoms (Phenotype)
X-linked			
Duchenne muscular dystrophy	310200	40–105	Muscle atrophy
Hemophilia A	306700	30–60	Severe impairment of blood clotting
Hemophilia B	306900	0.5–10	Mild impairment of blood clotting
Autosomal Dominant			
Achondroplasia	100800	10	Very short stature
Aniridia	106200	2.6	Absence of iris
Huntington disease	143100	<1	Uncontrollable movements, personality changes
Marfan syndrome	154700	4–6	Long limbs, weakened blood vessel walls
Neurofibromatosis type 1	162200	40–100	Brown skin spots, benign tumors under skin
Osteogenesis imperfecta	166200	10	Easily broken bones
Polycystic kidney disease	600666	60–120	Benign growths in kidneys
Retinoblastoma	180200	5–12	Malignant tumor of retina

Mutational Hot Spots

In some genes mutations are more likely to occur in regions called hot spots, where sequences are repetitive. It is as if the molecules that guide and carry out replication become “confused” by short repeated sequences, much as an editor scanning a manuscript might miss the spelling errors in the words “happiness” and “bananana” (figure 12.6). For example, more than one-third of the many mutations that cause alkaptonuria occur at or near one or more CCC repeats, even though these repeats account for only 9 percent of the gene (see the opening essay for chapter 5).

The increased incidence of mutations in repeats has a physical basis. Within a gene, when DNA strands locally unwind to replicate in symmetrical or repeated sequences, bases located on the same strand may pair. A stretch of ATATAT might pair with TATATA elsewhere on the same strand, creating a loop that interferes with replication and repair enzymes, increasing risk of error. For example, mutations in the gene for clotting factor IX, which causes hemophilia B, occur 10 to 100 times as often at any of 11 sites in the gene that have extensive direct repeats of CG.

Small additions and deletions of DNA bases are more likely to occur near sequences called palindromes (see figure 12.6). These sequences read the same, in a 5' to 3' direction, on complementary strands. Put another way, the sequence on one strand is the reverse of the sequence on the complementary strand. Palindromes probably increase the spontaneous mutation rate by disturbing replication.

The blood disorder alpha thalassemia (MIM 141800) illustrates the confusing effect of direct (as opposed to inverted)

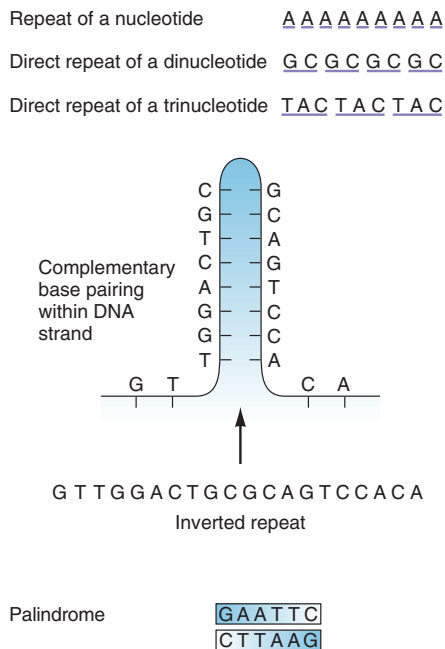


Figure 12.6 DNA symmetry may increase the likelihood of mutation. These examples show repetitive and symmetrical DNA sequences that may “confuse” replication enzymes, causing errors.

repeats of an entire gene. A person who does not have the disorder has four genes that specify alpha globin chains, two next to each other on each chromosome 16. Homologs with repeated genes can misalign during meiosis when the first sequence on one chromosome lies opposite the second sequence on the homolog. Crossing over can result in a sperm or oocyte that has one or three alpha globin genes instead of the normal two (figure 12.7). Fertilization with a normal gamete then results in a zygote with one extra or one missing alpha globin gene. At least three dozen conditions result from this unequal crossing over, including colorblindness (see Reading 6.1).

The number of alpha globin genes affects health. A person with only three alpha globin genes produces enough hemoglobin, and is a healthy carrier. Individuals with only two copies of the gene are mildly anemic and tire easily, and a person with a single alpha globin gene is severely anemic. A fetus lacking alpha globin genes does not survive.

Induced Mutation

Researchers can sometimes infer a gene’s normal function by observing what happens when mutation alters it. Because the spontaneous mutation rate is far too low to be a practical source

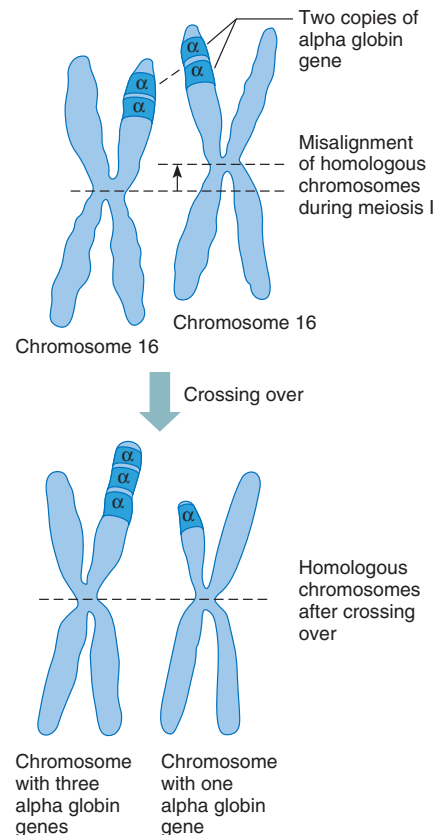


Figure 12.7 Gene duplication and deletion. The repeated alpha globin genes are prone to mutation by mispairing during meiosis.

of genetic variants for experiments, researchers make mutants. Geneticists use mutagens on model organisms to infer normal gene functions, yielding many collections and insights into human health.

Intentional Use of Mutagens

Chemicals or radiation are used to induce mutation. Alkylating agents, for example, are chemicals that remove a DNA base, which is replaced with any of the four bases—three of which are a mismatch against the complementary strand. Dyes called acridines add or remove a single DNA base. Because the DNA sequence is read three bases in a row, adding or deleting a single base can destroy a gene's information, altering the amino acid sequence of the encoded protein. Several other mutagenic chemicals alter base pairs, so that an A-T replaces a G-C, or vice versa. X rays and other forms of radiation delete a few bases or break chromosomes.

Researchers have developed several ways to test the mutagenicity of a substance. The best known, the Ames test, assesses how likely a substance is to harm the DNA of rapidly reproducing bacteria. One version of the test uses a strain of *Salmonella* that cannot grow when the amino acid histidine is absent from its medium. If exposure to a substance enables bacteria to grow on the deficient medium, then a gene has mutated that allows it to do so. A variation of the Ames test uses mammalian liver tissue.

In another variation of the Ames test, researchers exposed human connective tissue cells growing in culture to liquefied cigarette smoke. The chemicals from the smoke cut chromosomes through both DNA strands. This is an especially damaging insult because broken chromosomes can join with each other in different ways that can activate cancer-causing genes. Hence, the experiment may have modeled one way that cigarettes cause cancer.

Because many mutagens are also carcinogens (cancer-causing agents), the substances that the Ames test identifies as mutagens may also cause cancer. **Table 12.4** lists some common mutagens.

A limitation of using a mutagen is that it cannot cause a specific mutation. In contrast, a technique called site-directed mutagenesis changes a gene in a desired way. A gene is mass-produced, but it includes an intentionally substituted base, just as an error in a manuscript is printed in every copy of a book. Site-directed mutagenesis is faster and more precise than waiting for nature or a mutagen to produce a useful variant. It also makes it possible to study lethal mutations that can theoretically exist, but never do because they are so drastic that development does not proceed very far. Researchers can study such a lethal mutation in cell culture, or in model organisms such as flies or mice before they cease developing.

Accidental Exposures to Mutagens

Some mutagen exposure is unintentional. This occurs from workplace contact before the danger is known; from industrial accidents; from medical treatments such as chemotherapy and radiation; and from exposure to weapons that emit radiation.

An environmental disaster that released mutagenic radiation was a steam explosion at a nuclear reactor in the former Soviet Union on April 25, 1986. Between 1:23 and 1:24 A.M., Reactor 4 at the Chernobyl Nuclear Power Station in Ukraine exploded, sending a great plume of radioactive isotopes into the air that spread for thousands of miles. The reactor had been undergoing a test, its safety systems temporarily disabled, when it overloaded and rapidly flared out of control. Twenty-eight people died of acute radiation exposure in the days following the explosion.

Acute radiation poisoning is not genetic. Evidence of a mutagenic effect is the increased rate of thyroid cancer among children who were living in nearby Belarus. Rates have multiplied tenfold. The thyroid glands of young people soak up iodine, which in a radioactive form bathed Belarus in the days after the explosion. Cancer rates have also risen among workers who cleaned up the disaster. Analysis of radiation exposure in their teeth is being used to assess whether cancer risk rises with degree of exposure.

Another way researchers tracked mutation rates after the Chernobyl explosion was to compare the lengths of short DNA repeats, called minisatellite sequences, in children born in 1994 and in their parents, who lived in the Mogilev district of Belarus at the time of the accident and have remained there. Minisatellites are the same length in all cells of an individual. A minisatellite size in a child that does not match the size of either parent indicates that a mutation occurred in a parent's gamete. Such a mutation was twice as likely to occur in exposed families as in families living elsewhere. Mutation rates of nonrepeated DNA

Table 12.4 Commonly Encountered Mutagens

Mutagen	Source
Aflatoxin B	Fungi growing on peanuts and other foods
2-amino 5-nitrophenol	Hair dye components
2,4-diaminoanisole	"
2,5-diaminoanisole	"
2,4-diaminotoluene	"
p-phenylenediamine	"
Furylfuramide	Food additive
Nitrosamines	Pesticides, herbicides, cigarette smoke
Proflavine	Antiseptic in veterinary medicine
Sodium nitrite	Smoked meats
Tris (2,3-dibromopropyl phosphate)	Flame retardant in children's sleepwear

sequences are too low to provide useful information on the effects of radiation exposure, so investigators track minisatellites as a sensitive test of change.

Researchers learned of a new type of mutation from a young man conceived within a week of the Chernobyl accident, near the disaster site. He has extra digits, an abnormal epiglottis, and a benign growth on the hypothalamus, a group of symptoms called Pallister-Hall syndrome (MIM 146510). On his way to a camp for “children of Chernobyl” in the summer of 2002, he stopped at the National Institutes of Health to provide a DNA sample. Researchers indeed found a mutation in the gene on chromosome 7 known to cause the syndrome—a 72-base insertion that causes a “stop” codon to form, shortening the encoded protein. Oddly, the insertion matched mitochondrial DNA sequences. Apparently, the radiation damaged mitochondria in the sperm or oocyte, sending some mitochondrial DNA into the nucleus, where it inserted into the Pallister-Hall gene. Another clue to the unusual origin of this young man’s condition is that it is autosomal dominant, but neither of his parents have it. Since his case, researchers have discovered other such “nuclear DNA sequences of mitochondrial origin.”

Natural Exposure to Mutagens

Simply being alive exposes us to radiation that can cause mutation. Natural environmental sources of radiation include cosmic rays, sunlight, and radioactive minerals in the earth’s crust, such as radon. Contributions from medical X rays and occupational radiation hazards are comparatively minor (**table 12.5**). Job sites with increased radiation exposure include weapons facilities, research laboratories, health care facilities, nuclear power plants, and certain manufacturing plants (**figure 12.8**). Radiation exposure is measured in units called millirems; the average annual exposure in the northern hemisphere is 360 millirems.

Most of the potentially mutagenic radiation we are exposed to is of the ionizing type, which means that it has sufficient energy to remove electrons from atoms. Unstable atoms that emit ionizing radiation both exist naturally and are made by humans. Ionizing radiation breaks the DNA sugar-phosphate backbone.

Table 12.5 Sources of Radiation Exposure	
Source	Percentage of Total
Natural (cosmic rays, sunlight, earth’s crust)	81%
Medical X rays	11%
Nuclear medicine procedures	4%
Consumer products	3%
Other (nuclear fallout, occupational)	<1%

Ionizing radiation is of three major types. Alpha radiation is the least energetic and most short-lived, and the skin absorbs most of it. Uranium and radium emit alpha radiation. Beta radiation can penetrate the body farther, and emitters include tritium (a form of hydrogen), carbon-14, and strontium-70. Both alpha and beta rays tend not to harm health, although they can do damage if inhaled or eaten. In contrast is the third type of ionizing radiation, gamma rays. These can penetrate the body, damaging tissues. Plutonium and cesium isotopes used in weapons emit gamma rays, and this form of radiation is used to kill cancer cells.

X rays are the major source of exposure to human-made radiation, and they are not a form of ionizing radiation. They have less energy and do not penetrate the body to the extent that gamma rays do.

The effects of radiation damage to DNA depend upon the functions of the mutated genes. Mutations in oncogenes or tumor suppressor genes, discussed in chapter 18, can cause cancer. Radiation damage can be widespread, too. Exposing cells to radiation and then culturing them causes a genome-wide destabilization, so that mutations may occur even after the cell has divided a few times. Cell culture studies have also identified a “bystander effect,” when radiation harms cells not directly exposed.

Chemical mutagens are in the environment, too. Evaluating the risk that a specific chemical exposure will cause a mutation is very difficult, largely because people vary greatly in inherited susceptibilities, and are exposed to many chemicals. The risk that exposure to a certain chemical will cause a mutation is often less than the natural variability in susceptibility within a population, making it nearly impossible to track the true source and mechanism of any mutational event. Genetic tests can be used to determine specific inherited risks for specific employees who might encounter a mutagen in the workplace. However, in the United States, the Genetic Information Nondiscrimination Act (GINA) prevents employers from using such information to discriminate against employees.



Figure 12.8 Radiation and DNA. Exposure to radiation can cause mutation, whether from an unnatural source, such as this nuclear power plant, or a natural source, such as the sunlight that enables the field of rapeseed in the foreground to grow.

Key Concepts

1. Genes have different mutation rates.
2. Spontaneous mutations result when rare bases are incorporated during replication.
3. Spontaneous mutations are more frequent in viruses and microorganisms because they reproduce often.
4. Mutations are more likely in repetitive DNA.
5. Mutagens are chemicals or radiation that increase the risk of mutation. Researchers use mutagens to more quickly obtain mutants, which reveal normal gene function. Site-directed mutagenesis creates and amplifies specific mutations.
6. Mutagens are encountered in the natural environment and may be released in industrial accidents.

12.4 Types of Mutations

Mutations can be classified by whether they remove, alter, or add a function, or by exactly how they structurally alter DNA. The same single-gene disorder can result from different types of mutations, as **figure 12.9** shows for familial hypercholesterolemia. **Table 12.6** summarizes the types of mutations described in this section using an analogy to an English sentence.

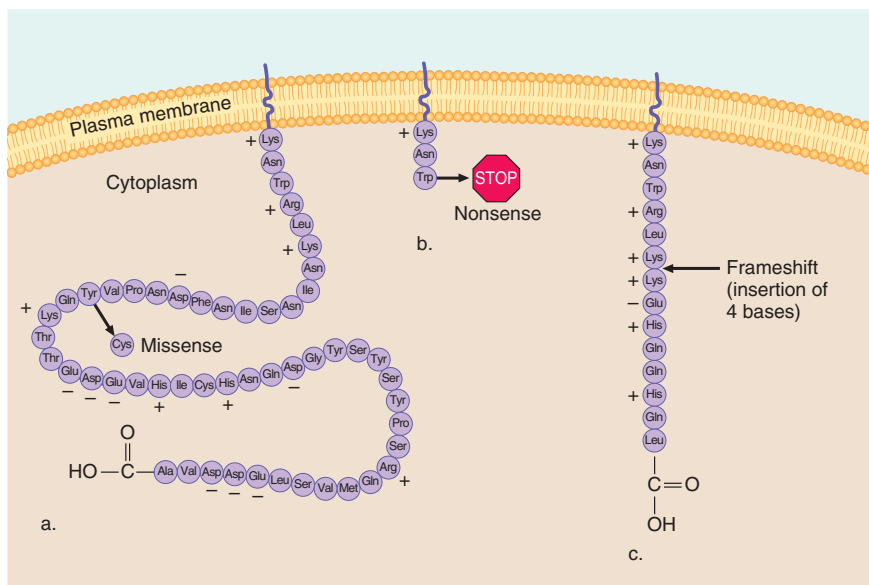


Figure 12.9 Different mutations in a gene can cause the same disorder. In familial hypercholesterolemia, several types of mutations alter the LDL receptor normally anchored in the plasma membrane. LDL receptor **(a)** bears a missense mutation—a cysteine substitutes for a tyrosine bending the receptor enough to impair its function. The short LDL receptor in **(b)** results from a nonsense mutation, in which a stop codon replaces a tryptophan codon. In **(c)**, a 4-base insertion alters the reading frame.

Point Mutations

A **point mutation** is a change in a single DNA base. It is a **transition** if a purine replaces a purine (A to G or G to A) or a pyrimidine replaces a pyrimidine (C to T or T to C). It is a **transversion** if a purine replaces a pyrimidine or vice versa (A or G to T or C). A point mutation can have any of several consequences—or it may have no obvious effect at all on the phenotype, acting as a silent mutation.

Missense and Nonsense Mutations

A point mutation that changes a codon that normally specifies a particular amino acid into one that codes for a different amino acid is called a **missense mutation**. If the substituted amino acid alters the protein's conformation significantly or occurs at a site critical to its function, signs or symptoms of disease or an observable variant of a trait may result. About a third of missense mutations harm health.

The point mutation that causes sickle cell disease (see figure 12.2) is a missense mutation. The DNA sequence CTC encodes the mRNA codon GAG, which specifies glutamic acid. In sickle cell disease, the mutation changes the DNA sequence to CAC, which encodes GUG in the mRNA, which specifies valine. This mutation changes the protein's shape, which alters its function.

A point mutation that changes a codon specifying an amino acid into a “stop” codon—UAA, UAG, or UGA in mRNA—is a **nonsense mutation**. A premature stop codon shortens the protein product, which can greatly influence the phenotype.

For example, in factor XI deficiency (MIM 264900), a blood clotting disorder, a GAA codon specifying glutamic acid is changed to UAA, signifying “stop.” The shortened clotting factor cannot halt the profuse bleeding that occurs during surgery or from injury. Nonsense mutations are predictable by considering which codons can mutate to a “stop” codon.

In the opposite of a nonsense mutation, a normal stop codon mutates into a codon that specifies an amino acid. The resulting protein is longer than normal, because translation continues through what is normally a stop codon.

Point mutations can control transcription, affecting the quantity rather than the quality of a protein. For example, in 15 percent of people who have Becker muscular dystrophy (MIM 310200)—a milder adult form of the condition—the muscle protein dystrophin is normal, but its levels are reduced. The mutation is in the promoter for the dystrophin gene. This

Table 12.6 Types of Mutations

A sentence comprised of three-letter words can provide an analogy to the effect of mutations on a gene's DNA sequence:

Normal	THE ONE BIG FLY HAD ONE RED EYE
Missense	THQ ONE BIG FLY HAD ONE RED EYE
Nonsense	THE ONE BIG [REDACTED]
Frameshift	THE ONE QBI GFL YHA DON ERE DEY
Deletion	THE ONE BIG [REDACTED] HAD ONE RED EYE
Insertion	THE ONE BIG WET FLY HAD ONE RED EYE
Duplication	THE ONE BIG FLY FLY HAD ONE RED EYE
Expanding mutation	
generation 1	THE ONE BIG FLY HAD ONE RED EYE
generation 2	THE ONE BIG FLY FLY FLY HAD ONE RED EYE
generation 3	THE ONE BIG FLY FLY FLY FLY FLY HAD ONE RED EYE

slows transcription, and dystrophin protein is scarce. Muscle function suffers. The other 85 percent of individuals who have Becker muscular dystrophy have shortened proteins, not a deficiency of normal-length proteins.

Another way that point mutations can affect protein production is to disrupt the trimming of long precursor molecules. Such a mutation causes the type of Ehlers-Danlos syndrome that affects the boy shown in figure 12.4.

Splice Site Mutations

A point mutation can greatly affect a gene's product if it alters a site where introns are normally removed from the mRNA. This is called a splice site mutation. It can affect the phenotype if an intron is translated into amino acids, or if an exon is skipped instead of being translated, shortening the protein.

Retaining an intron is unusual because most introns have stop codons in all reading frames. However, if a stop codon is not encountered, a retained intron adds bases to the protein-coding portion of an mRNA. For example, in one family with severe cystic fibrosis, a missense mutation alters an intron site so that it is not removed. The encoded protein is too bulky to move to its normal position in the plasma membrane.

A missense mutation can cause "exon skipping" that need not alter the amino acid sequence, but removes a few amino acids. The mutation creates an intron splicing site where there should not be one, and an entire exon is "skipped" when the mRNA is translated into protein, as if it were an intron. An exon-skipping mutation is a deletion at the mRNA level, but a missense mutation at the DNA level. For example, a disorder called familial dysautonomia (MIM 223900) (FD) usually results from exon skipping in the gene encoding an enzyme necessary for the survival of certain neurons that control

sensation and involuntary responses. The *In Their Own Words* box on page 226 describes life for a child with FD.

A peculiarity of some disorders caused by exon skipping is that some cells ignore the mutation and manufacture a normal protein from the affected gene—after all, the amino acid sequence information is still there. Depending upon which cells actually make the encoded protein, the phenotype may be less severe than in individuals with the same disorder but with a different type of mutation in an exon.

Studies on various cell types from individuals who have FD or have died from the disease reveal that the cells in which the exon is skipped are the cells that contribute to symptoms. That is, many cells from the brain and spinal cord skip the exon, but cells from muscle, lung, liver, white blood cells, and various glands produce normal-length proteins. This means there may be a way to coax nervous system cells in affected children to also produce the protein. Current clinical trials are examining the ability of several natural compounds to restore normal processing of the

FD gene's information.

Deletions and Insertions

In genes, the number 3 is very important, because triplets of DNA bases specify amino acids. Adding or deleting a number of bases that is not a multiple of three devastates a gene's function because it disrupts the gene's reading frame, which refers to the nucleotide position where the DNA begins to encode protein (see figure 10.14). An exon is usually "readable" (has no stop codons) in only one of its three possible reading frames. A change that alters the reading frame is called a **frameshift mutation**. Line 4 in table 12.6 illustrates a frameshift mutation.

A **deletion mutation** removes genetic material. A deletion that removes three or a multiple of three bases will not cause a frameshift, but can still alter the phenotype. Deletions range from a single DNA nucleotide to thousands of bases to large parts of chromosomes. Chapter 13 considers large deletions. Many common inherited disorders result from deletions, including male infertility caused by tiny deletions in the Y chromosome.

An **insertion mutation** adds DNA and it, too, can offset a gene's reading frame. In one form of Gaucher disease, for example, an inserted single DNA base prevents production of an enzyme that normally breaks down glycolipids in lysosomes. The resulting buildup of glycolipid enlarges the liver and spleen and causes easily fractured bones and neurological impairment.

Another type of insertion mutation repeats part of a gene's sequence. The insertion is usually adjacent or close to the original sequence, like a typographical error repeating a word word. Two copies of a gene next to each other

In Their Own Words

Familial Dysautonomia: Rebekah's Story

Our daughter Rebekah has familial dysautonomia. This is a rare genetic disorder that affects the autonomic and peripheral nervous systems. Rebekah was born in 1992, but she was not diagnosed until she was almost 3. She appeared to be healthy at birth, but she began to decline rapidly by 9 months. Rebekah suffered from frequent pneumonia, vomiting and retching, extremely high fevers, chills, rapid heartbeat, and seizures. At times, she would become covered with hot, red blotches. Other times, her hands and feet got very cold and appeared puffy and blue. Episodes of crying would precipitate breathholding, when she would turn blue and lose consciousness. As she lost ground on the growth and development charts, medical testing failed to reveal a cause for these symptoms. We wondered if we would identify the problem before she died. Our physicians sometimes hinted that perhaps we were the cause.

After more than twelve local hospitalizations and a variety of tests, we traveled to a major children's teaching hospital, hoping that a fresh team of doctors would identify Rebekah's condition. One doctor knew immediately that she had FD. He recognized the pattern of "dysautonomic crises." Two more symptoms, which we hadn't even noticed, were diagnostic indicators. Individuals with FD do not cry tears, and they lack papillae (bumps) on the tip of the tongue. Our Eastern European, Jewish heritage was also a clue, because FD is one of a number of diseases primarily affecting this population.

To a varying degree, FD reduces sensation of pain, heat, and cold. There are problems with balance and coordination, including motor difficulties that affect feeding, swallowing, and breathing. Most people with FD have a feeding tube, and must limit what they eat or drink by mouth due to danger of aspiration. FD causes fluctuations in blood pressure, digestive problems, and learning disabilities. Most individuals develop scoliosis, usually requiring corrective spine surgery. In short, FD affects every organ and system in the body.

The diagnosis allowed us to finetune Rebekah's therapies and activities to maximize her well-being. With improved nutrition, excellent therapies, and wonderful teachers, Rebekah has made tremendous progress, but we are always poised for a hospital stay. Even a minor illness can set off a crisis. A team of pediatric specialists monitors her lungs, heart, eyes, back, and growth and development.

Rebekah is a happy, good-natured child who makes friends easily and is sensitive to the needs of others. She works hard in school, and is able to keep up when her health is good. She has learned to overcome her learning challenges, using assistive technology in school to help her with writing and organizing. When Rebekah got a back brace to try to slow the scoliosis, and I tried to steer her to choose clothes that would deemphasize the bulk of the brace, she told me, "Mom, just relax. They're going to see it sooner or later!"

We don't know what the future holds. FD is a progressive, degenerative disease with life-threatening complications and a shortened lifespan. Any major stress, including developmental changes, surgery, a serious illness, and increased emotional stress, can exacerbate the severity. Yet, we feel hopeful for our daughter's future. We are most encouraged by Rebekah herself. Her positive outlook on life, her willingness to find the good in any situation, and her ability to overcome challenges with spunk and humor inspire everyone around her.

Lynn Lieberman



Rebekah with her dog, Tracy.

is called a **tandem duplication**. A form of Charcot-Marie-Tooth disease (MIM 118200), which causes numb hands and feet, results from a one-and-a-half-million-base-long tandem duplication.

Figure 12.9 compares the effects on protein sequence of missense, nonsense, and frameshift mutations in the gene that encodes the LDL receptor, causing familial hypercholesterolemia (see figure 5.2). These three mutations exert very different effects on the protein. A missense mutation replaces one amino acid with another, bending the protein in a way that

impairs its function. A nonsense mutation greatly shortens the protein. A frameshift mutation introduces a section of amino acids that is not in the wild type protein.

Pseudogenes and Transposons Revisited

Recall from chapter 11 that a pseudogene is a DNA sequence that is very similar to the sequence of a protein-encoding gene. A pseudogene is not translated into protein, although it may be transcribed. The pseudogene may have descended from the

original gene sequence, which was duplicated when DNA strands misaligned during meiosis, similar to the situation depicted in figure 12.7 for the alpha globin gene. When this happens, a gene and its copy end up right next to each other on the chromosome. The original gene or the copy then mutates to such an extent that it is no longer functional and becomes a pseudogene. Its duplicate lives on as the functional gene.

Although a pseudogene is not translated, its presence can interfere with the expression of the functional gene and cause a mutation. For example, some cases of Gaucher disease result from a crossover between the working gene and its pseudogene, which has 96 percent of the same sequence and is located 16,000 bases away. The result is a fusion gene, which is a DNA sequence containing part of the functional gene and part of the pseudogene. The fusion gene does not retain enough of the normal gene sequence to enable the cell to synthesize the encoded enzyme, and Gaucher disease results. Gaucher disease is a lysosomal storage disease that causes fatigue, bruising, anemia, and weak bones. The phenotype is very variable, and for many patients, supplying the enzyme eliminates symptoms.

Chapter 11 also considered transposons, or “jumping genes.” Transposons can alter gene function in several ways. They can disrupt the site they jump from, shut off transcription of the gene they jump into, or alter the reading frame of their destination if they are not a multiple of three bases. For example, a boy with X-linked hemophilia A had a transposon in his factor VIII gene—a sequence that was also in his carrier mother’s genome, but on her chromosome 22. Apparently, in the oocyte, the transposon jumped into the factor VIII gene on the X chromosome, causing the boy’s hemophilia.

Expanding Repeats

Myotonic dystrophy is an inherited disease that begins at an earlier age and causes more severe symptoms from one generation to the next. A grandfather might experience only mild weakness in his forearms, and cataracts. His daughter might have more noticeable arm and leg weakness, and a flat facial expression. Her affected children might experience severe muscle weakness.

For many years, the “anticipation”—the worsening of symptoms over generations—was thought to be psychological. Then, with the ability to sequence genes, researchers found that myotonic dystrophy indeed worsens with each generation because the gene expands! Myotonic dystrophy is caused by a type of mutation called an expanding triplet repeat. The gene, on chromosome 19, has an area rich in repeats of the DNA triplet CTG. A person who does not have myotonic dystrophy usually has from 5 to 37 copies of the repeat; a person with the disorder has from 50 to thousands of copies (figure 12.10).

Expanding triplet repeats have been discovered in more than fifteen human inherited disorders. Usually, a repeat number of fewer than 40 copies is stably transmitted to the next generation and doesn’t produce symptoms. Larger repeats are unstable, enlarging with each generation and causing symptoms

Pedigree		Age of onset	Phenotype	Number of copies of GAC mRNA repeat
I		Older adulthood	Mild forearm weakness, cataracts	50–80
II		Mid-adulthood	Moderate limb weakness	80–700
III		Childhood	Severe muscle impairment, respiratory distress, early death	700+

Figure 12.10 Expanding genes explain anticipation. In some disorders, symptoms that worsen from one generation to the next—termed *anticipation*—have a physical basis: The gene is expanding as the number of repeats grows.

that are more severe and begin sooner. **Reading 12.1** describes the triplet repeat disorder fragile X syndrome.

The mechanism behind triplet repeat disorders lies in the DNA sequence. The bases of the repeated triplets implicated in the expansion diseases, unlike others, bond to each other in ways that bend the DNA strand into shapes, such as hairpins. These shapes then interfere with replication, which causes the expansion. Once these repeats are translated, the extra-long proteins shut down cells in various ways:

- binding to parts of transcription factors that have stretches of amino acid repeats similar to or matching the expanded repeat
- blocking proteasomes and thereby enabling misfolded proteins to persist
- directly triggering apoptosis

Triplet repeat proteins may also enter the nucleus, even though their wild type versions function only in the cytoplasm, or vice versa.

The triplet repeat disorders are said to cause a “dominant toxic gain of function.” This means that they cause something novel to happen, rather than removing a function, such as is often associated with recessive enzyme deficiencies. The idea of a gain of function arose from the observation that deletions of these genes do not cause symptoms.

Table 12.7 describes several triplet repeat disorders. Particularly common among them are the “polyglutamine diseases” that have repeats of the mRNA codon CAG, which encodes the amino acid glutamine.

For some triplet repeat disorders, the mutation thwarts gene expression before a protein is even manufactured. In myotonic dystrophy type 1 the expansion is in the initial untranslated region of a gene on chromosome 19, resulting in a huge mRNA. When genetic testing became available for the disorder, researchers discovered a second form of the illness in people who had wild type alleles for the

Reading 12.1

Fragile X Mutations Affect Boys and Their Grandfathers

In the 1940s, geneticists hypothesized that a gene on the X chromosome confers mental retardation, because more affected individuals are male. It wasn't until 1969, though, that a clue emerged to the genetic basis of X-linked mental retardation. Two retarded brothers and their mother had an unusual X chromosome. The tips at one chromosome end dangled, separated from the rest of each chromatid by a thin thread (**figure 1a**). When grown under specific culture conditions (lacking folic acid), this part of the X chromosome was very prone to breaking—hence, the name fragile X syndrome. Worldwide, it affects 1 in 2,000 males, accounting for 4 to 8 percent of all males with mental retardation. One in 4,000 females is affected. They usually have milder cases because of the presence of a second, normal X chromosome.

Youngsters with fragile X syndrome look normal, but by young adulthood, their faces are very long and narrow, with a long jaw and protruding ears. The testicles are very large. Mental impairment and behavioral problems include mental retardation, learning disabilities, repetitive speech, hyperactivity, shyness, social anxiety, a short attention span, language delays, and temper outbursts.

Fragile X syndrome is inherited in an unusual pattern. Because the fragile chromosome is the X, the associated syndrome should be transmitted as any X-linked trait is, from carrier mother to affected son. However, penetrance is incomplete. One-fifth of males who inherit the chromosomal abnormality have no symptoms. But because they pass on the affected chromosome to all their daughters—half of whom have some degree of mental impairment—they are called “transmitting males.” A transmitting male's grandchildren may inherit fragile X syndrome.

A triplet repeat mutation causes fragile X syndrome. In unaffected individuals, the fragile X area contains about 30 repeats of the sequence CGG, in a gene called the fragile X mental retardation gene (*FMR1*). In people who have the fragile chromosome and show its effects, this region is expanded to 200 to 2,000 CGG repeats. Transmitting males, as well as females with mild symptoms, or who have affected sons, often have a premutation consisting of 50 to 200 repeats.

The *FMR1* gene encodes fragile X mental retardation protein (FMRP). This protein, when abnormal, binds to and disables several different mRNA molecules whose encoded proteins are crucial for brain neuron function.

Despite our knowledge of the genetic mechanism behind fragile X syndrome, mysteries remain. A distinct type of disorder has been described in the maternal grandfathers of boys who have fragile X syndrome. Clinicians noticed that mothers of boys with fragile X syndrome very often reported the same symptoms in their fathers—tremors, balance problems, and then cognitive or psychiatric difficulties (inability to plan or pay attention,

and inappropriate behaviors). The grandfathers were sometimes misdiagnosed with Parkinson disease due to the tremors. However, Parkinson's patients can pass the “tandem gait test” (walking a straight line) while the grandfathers could not. The grandfathers' symptoms worsen with time and can lead to premature death (**table 1**).

Further investigation led to the description of the new condition, called fragile X-associated tremor/ataxia syndrome (FXTAS, MIM 300623). (Ataxia is poor balance and coordination.) The disorder has been studied in brains obtained after the grandfathers died and in a mouse model. Like the granddads, the mice are fine until middle age. Then they, too, develop tremors and balance problems as well as nervousness and memory impairment. Perhaps the symptoms of FXTAS arise from excess *FMR1* mRNA, which attracts and disables other mRNAs.

The discovery of FXTAS has genetic counseling implications. As neurologists learn to distinguish this disorder from others, such as Parkinson disease, daughters can be counseled that they might pass on the condition to sons, and be offered testing.

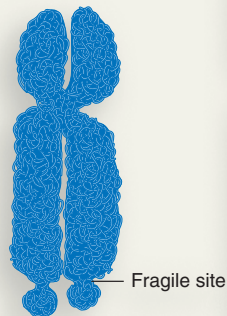
Table 1

Prevalence of FXTAS in
Grandfathers of Fragile
X Syndrome Grandsons

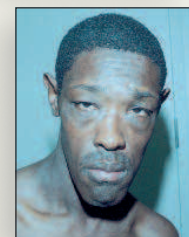
Age	Prevalence
50s	17%
60s	38%
80+	75%



a.



Fragile site



b.

Figure 1 Fragile X syndrome. A fragile site on the tip of the long arm of the X chromosome (**a**) is associated with mental retardation and a characteristic long face that becomes pronounced with age (**b**).

Table 12.7 Triplet Repeat Disorders

Disorder	MIM	Repeat	Normal Copy #	Disease Copy #	Signs and Symptoms (Phenotype)
Fragile X syndrome	309550	CGG or CCG	6–50	200–2,000	Mental retardation, large testicles, long face
Friedreich ataxia	229300	GAA	6–29	200–900	Loss of coordination and certain reflexes, spine curvature, knee and ankle jerks
Haw River syndrome	140340	CAG	7–25	49–75	Loss of coordination, uncontrollable movements, dementia
Huntington disease	143100	CAG	10–34	40–121	Personality changes, uncontrollable movements, dementia
Huntington disease-like 2	606438	CAG	8–28	40–58	Identical to Huntington disease
Jacobsen syndrome	147791	CGG	11	100–1,000	Poor growth, abnormal face, slow movement
Myotonic dystrophy type I	160900	CTG	5–37	80–1,000	Progressive muscle weakness; heart, brain, and hormone abnormalities
Myotonic dystrophy type II	602668	CCTG	<10	>100	Progressive muscle weakness; heart, brain, and hormone abnormalities
Spinal and bulbar muscular atrophy	313200	CAG	14–32	40–55	Muscle weakness and wasting in adulthood
Spinocerebellar ataxia (7 types)	271245	CAG	4–44	40–130	Loss of coordination

chromosome 19 gene. They have myotonic dystrophy type 2, which is caused by an expanding *quadruple* repeat of CCTG in a gene on chromosome 3. Affected individuals have more than 100 copies of the repeat, compared to the normal fewer than 10 copies.

When researchers realized that this second repeat mutation for myotonic dystrophy was also in a non-protein-encoding part of the gene—an intron—a mechanism of disease became apparent: The mRNA is not processed normally and as a result cannot exit the nucleus. In myotonic dystrophy type 1, the excess material is added to the start of the gene; in type 2, it appears in an intron that is not excised. The bulky mRNAs bind to a protein that, in turn, alters intron splicing in several other genes. Deficiency of the proteins encoded by these final affected genes causes the symptoms.

A lesson learned from the expanding repeat disorders is that a DNA sequence is more than just one language that can be translated into another. Whether a sequence is random—CGT CGT ATG CAT CAG, for example—or highly repetitive—such as CAG CAG CAG CAG and on and on—can affect transcription, translation, or the ways that proteins interact.

Copy Number Variants

In addition to differing, if only slightly, in our DNA sequences, we also differ in the numbers of copies of particular DNA sequences. These sequences that vary in number from person to person are called **copy number variants** (CNVs). Our genomes have hundreds to thousands of them, and they

account for about a quarter of the genome. Researchers were not aware of the extent and abundance of CNVs until recently, because the technology used to sequence the human genome detected any DNA sequence only once—a little like searching this book for the word “variant,” and not the number of times it is used.

Copy number is a different form of information than DNA sequence differences. A language metaphor is useful to distinguish point mutations and single nucleotide polymorphisms (SNPs) from CNVs. If a wild type short sequence and a variant with two SNPs are written as:

The fat rat sat on a red cat (wild type)

The fat rat sat in a red hat (two SNPs)

then the sequence with two CNVs might be:

The fat fat rat sat on a red red red cat

CNVs may contribute significantly to the differences among us. A CNV can range in size from a few DNA bases to millions, and copies may lie next to each other on a chromosome (“tandem”) or might be far away—even parts of other chromosomes.

CNVs may have no effect on the phenotype, or they can disrupt a gene’s function and harm health. A CNV may have a direct effect by inserting into a protein-encoding gene and offsetting its reading frame, or have an indirect effect by destabilizing surrounding sequences.

The first CNV to be found in a family with a single-gene disorder consists of about 750,000 bases repeated in tandem

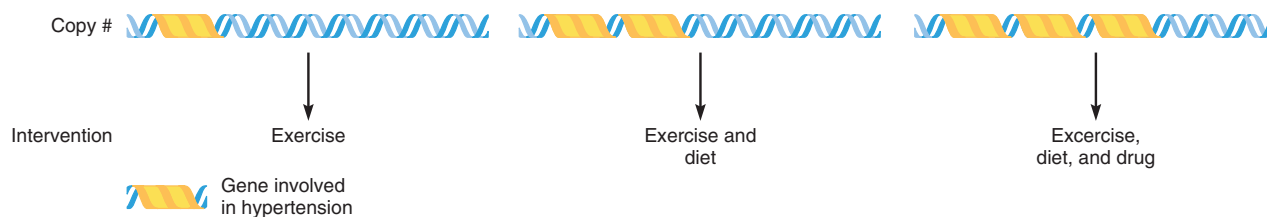


Figure 12.11 Using copy number variants in health care. If large-scale studies support a correlation of copy number of a particular DNA sequence with cholesterol level, then perhaps the number can be used to give medical advice: exercise, exercise plus dietary changes, exercise plus diet plus a cholesterol-lowering drug.

five times, at the tip of the short arm of chromosome 4. It causes an autosomal dominant syndrome in a large family in Belgium. Affected family members have very small ears, holes in the eyes, and blocked tear ducts.

Detecting CNVs may be useful in health care even if we do not yet know exactly what they do, if specific CNVs correlate to specific phenotypes in many people. Such an approach is similar to genome-wide association studies based on SNPs—the more people examined, the more powerful the connection. **Figure 12.11** depicts, schematically, how CNVs correlated to cholesterol level might be used to guide medical advice.

We still have much to learn about copy number variants. It isn't yet known how many are correlated to phenotypes (disorders), and the range of such repeats in what we can call a “normal” human genome.

Key Concepts

1. A point mutation alters a single DNA base and can occur in any part of a gene.
2. In a transversion, a purine replaces a pyrimidine, or vice versa; in a transition, a purine replaces a purine or a pyrimidine replaces a pyrimidine.
3. A missense mutation replaces one amino acid with another.
4. A nonsense mutation changes a coding codon into a “stop” codon, shortening the protein. A stop codon that is changed to a coding codon lengthens the protein.
5. Mutations in intron/exon splice sites, promoters, or other control regions affect gene function.
6. Inserting or deleting bases can shift the reading frame.
7. Tandem duplications repeat a DNA sequence.
8. Pseudogenes are nonfunctional sequences very similar to nearby functional genes.
9. Transposons can move, insert into genes, and cause illness.
10. Expanded repeats exert effects that arise from protein misfolding.
11. Copy number variants can affect the phenotype and distinguish individuals.

12.5 The Importance of Position

The degree to which a mutation alters the phenotype depends upon where in the gene the change occurs, and how the mutation affects the conformation, activity, or abundance of an encoded protein. A mutation that replaces an amino acid with a very similar one would probably not affect the phenotype greatly, because it wouldn't substantially change the conformation of the protein. Even substituting a very different amino acid would not have much effect if the change is in part of the protein not crucial to its function.

The effects of specific mutations are well-studied in hemoglobin. They are less understood, but still fascinating, in the gene that encodes prion protein.

Globin Variants

Because the globin gene mutations were the first to be analyzed in humans, and because some variants are easily detected using electrophoresis, hundreds of globin gene mutations have been known for years. Mutations in these genes can cause anemia with or without sickling, or cause cyanosis (a blue pallor due to poor oxygen binding). Rarely, a mutation boosts the molecule's affinity for oxygen. Some globin gene variants exert no effect and are thus considered “clinically silent” (**table 12.8**).

Oddly, hemoglobin S and hemoglobin C are variants that result from mutations that change the sixth amino acid in the beta globin polypeptide, but in different ways. Homozygotes for hemoglobin S have sickle cell disease, yet homozygotes for hemoglobin C are healthy. Both types of homozygotes are resistant to malaria because the unusual hemoglobin alters the shapes and surfaces of red blood cells in ways that keep out the parasite that causes the illness. The connection between sickle cell disease and malaria is discussed in chapter 15.

An interesting consequence of certain mutations in either the alpha or beta globin chains is hemoglobin M. Normally, the iron in hemoglobin is in the ferrous form, which means that it has two positive charges. In hemoglobin M, the mutation stabilizes the ferric form, which has three positive charges and cannot bind oxygen. Fortunately, an enzyme converts the abnormal ferric iron to the normal ferrous form, so that the only symptom is usually cyanosis. The condition has

Table 12.8 Globin Mutations

Associated Phenotype	Name	Mutation
Clinically silent	Hb Wayne	Single-base deletion in alpha gene causes frameshift, changing amino acids 139–141 and adding amino acids
	Hb Grady	Nine extra bases add three amino acids between amino acids 118 and 119 of alpha chain
Oxygen binding	Hb Chesapeake	Change from arginine to leucine at amino acid 92 of beta chain
	Hb McKees Rocks	Change from tyrosine to STOP codon at amino acid 145 in beta chain
Anemia	Hb Constant Spring	Change from STOP codon to glutamine elongates alpha chain
	Hb S	Change from glutamic acid to valine at amino acid 6 in beta chain causes sickling
	Hb Leiden	Amino acid 6 deleted from beta chain
Protection against malaria	Hb C	Change from glutamic acid to lysine at amino acid 6 in beta chain causes sickling

been known for more than two hundred years in a small town in Japan. Many people there have “blackmouth” because of the cyanosis caused by the faulty hemoglobin. The condition is autosomal dominant.

Even more noticeable than people with blackmouth are the “blue people of Troublesome Creek.” Seven generations ago, in 1820, a French orphan named Martin Fugate who settled in this area of Kentucky brought in a recessive gene that causes a form of methemoglobinemia. He was missing an enzyme (cytochrome b5 reductase) that normally catalyzes a reaction that converts a type of hemoglobin with poor oxygen affinity, called methemoglobin, back into normal hemoglobin by adding an electron. Martin’s wife was a carrier for this very rare disease. After extensive inbreeding in the isolated community—their son married his aunt, for example—a large pedigree of “blue people” of both sexes arose.

In “blue person disease,” the excess oxygen-poor hemoglobin causes a dark blue complexion. Carriers may have frighteningly bluish lips and fingernails at birth, which usually improve. This form of methemoglobinemia also affects the Navajo and Eskimos. Treatment is simple: A tablet of methylene blue, a commonly used dye, adds the electron back to methemoglobin, converting it to normal hemoglobin.

Susceptibility to Prion Disorders

For the prion protein gene, as with the globin genes, certain mutations exert extreme effects, while others do not. Recall from chapter 10 that a prion is a protein that assumes both stable and infectious conformations. A prion disease can be inherited, such as fatal familial insomnia, or acquired, such as developing variant Creutzfeldt-Jakob disease from eating beef from a cow that had bovine spongiform encephalopathy (“mad cow disease”). The prion protein has at least eight distinct conformations. The normal form of the protein has a central core

made up of helices. In a disease-causing form, the helices open into a sheet.

Precise genetic changes control the plasticity of the prion protein. The 129th amino acid is particularly important. In people who inherit prion disorders, amino acid 129 is either valine in all copies of the protein (genotype VV, causing the insomnia) or methionine in all copies (genotype MM, causing a form of Creutzfeldt-Jakob disease). These people are homozygous for this small part of the gene. Most people, however, are heterozygous, with valine in some prion proteins and methionine in others (genotype VM). Perhaps having two different amino acids at this position enables the proteins to assemble and to carry out their normal functions without damaging the brain.

A mutation at a different site in the prion protein gene raises the risk of brain disease even higher. Normally prion protein folds so that amino acid 129 is near amino acid 178, which is aspartic acid. People who inherit prion diseases are homozygous for the gene at position 129, and have another mutation that changes amino acid 178 to asparagine.

Factors That Lessen the Effects of Mutation

Mutation is a natural consequence of DNA’s ability to change. This flexibility is essential for evolution because it generates new variants, some of which may resist environmental change and enable a population or even a species to survive. However, many factors minimize the deleterious effects of mutations on phenotypes.

The genetic code protects against mutation. Recall from chapter 10 that synonymous codons specify the same amino acid. Mutation in the third codon position is effectively “silent” because the two codons are synonymous. For example, a change from RNA codon CAA to CAG does not alter the designated amino acid, glutamine, so a protein whose gene contains the change would not change. A change from one codon to a nonsynonymous one could affect the phenotype.

Other genetic code nuances prevent synthesis of very altered proteins. For example, mutations in the second codon position sometimes replace one amino acid with another that has a similar conformation, minimizing disruption of the protein's conformation. GCC mutated to GGC, for instance, replaces alanine with equally small glycine.

A **conditional mutation** affects the phenotype only under certain conditions. This can be protective if an individual avoids the exposures that trigger symptoms. Consider a common variant of the X-linked gene that encodes glucose 6-phosphate dehydrogenase (G6PD), an enzyme that immature red blood cells use to extract energy from glucose. One hundred million people worldwide have G6PD deficiency (MIM 305900), which can cause life-threatening hemolytic anemia, but only under rather unusual conditions—eating fava beans or taking certain antimalarial drugs (**figure 12.12**).

In the fifth century B.C., the Greek mathematician Pythagoras wouldn't allow his followers to consume fava beans—he had discovered that it would sicken some of them. During the Second World War, several soldiers taking the antimalarial drug primaquine developed hemolytic anemia. A study began shortly after the war to investigate the effects of the drug on volunteers at the Stateville Penitentiary in Joliet, Illinois. Researchers identified abnormal G6PD in people who developed anemia when they took the drug.

What do fava beans, antimalarial drugs, and dozens of other triggering substances have in common? They “stress” red blood cells by exposing them to oxidants, chemicals that strip electrons from other compounds. Without the enzyme, the stress bursts the red blood cells.

Another protection against mutation occurs in stem cells. When a stem cell divides to yield another stem cell and a progenitor or differentiated cell, the oldest DNA strands segregate with the stem cell, and the most recently replicated

DNA strands go to the more specialized daughter cells. This makes sense in organs where stem cells very actively yield specialized daughter cells, such as the skin and small intestine. Because mutations occur when DNA replicates, this skewed distribution of chromosomes sends the DNA most likely to harbor mutations into cells that will soon be shed (from a towel rubbed on skin or in a bowel movement) while keeping mutations away from the stem cells that must continually regenerate the tissues.

Key Concepts

1. How a mutation alters the phenotype depends upon its location in the gene.
2. Mutations in globin genes have a variety of effects. Mutations in two parts of the prion protein gene predispose to developing a prion disorder.
3. Genetic code degeneracy ensures that some third-codon-position mutations do not alter the specified amino acid. Changes in the second codon position often substitute a structurally similar amino acid.
4. Conditional mutations are expressed only in certain environments.
5. Preferential segregation of the oldest DNA strands to stem cells rather than daughter cells protects against mutation.

12.6 DNA Repair

Any manufacturing facility tests a product in several ways to see whether it has been assembled correctly. Mistakes in production are rectified before the item goes on the market—most of the time. The same is true for a cell's manufacture of DNA.

Damage to DNA becomes important when the genetic material is replicated, because the error in sequence is passed on to daughter cells. In response to damage, the cell may die by apoptosis or it may repair the error. If the cell doesn't die or the error is not repaired, cancer may result. Fortunately, DNA replication is incredibly accurate—only 1 in 100 million or so bases is incorrectly incorporated. This is quite an accomplishment, because DNA replicates approximately 10^{16} times during an average human lifetime. However, most such mutations occur in somatic cells, and do not affect the phenotype.

DNA polymerase as well as DNA damage response genes oversee the accuracy of replication. DNA repair consists of a cell's detecting damage and signaling systems in the cell that respond with death or repair. More than 50 DNA damage response genes have been identified.

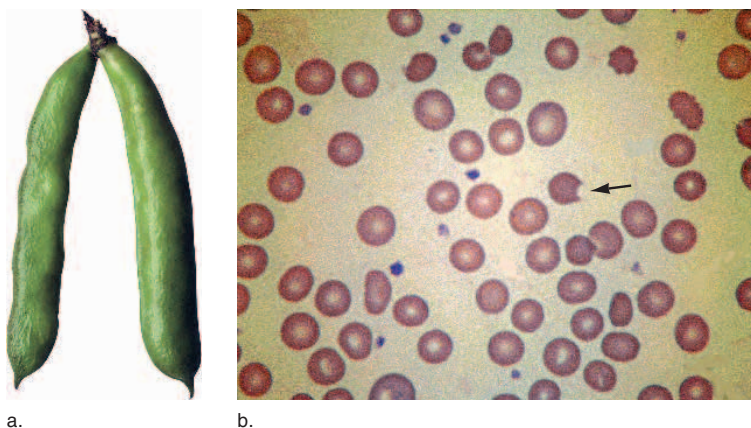


Figure 12.12 Sickness and circumstance. A conditional mutation causes some cases of G6PD deficiency hemolytic anemia. Exposure to two biochemicals in fava beans (**a**) causes hemoglobin molecules to unfold, which causes red blood cells to morph out of shape and then burst (**b**).

All eukaryotes can repair their nuclear DNA, although some species do so more efficiently than others. Mitochondrial DNA cannot repair itself, which accounts for its higher mutation rate. The master at DNA repair is a large, reddish microbe. *Deinococcus radiodurans* was discovered in a can of spoiled ground meat at the Oregon Agricultural Experiment Station in Corvallis in 1956, where it had withstood radiation used to sterilize the food. It tolerates 1,000 times the radiation level that a person can, and it can even live amidst the intense radiation of a nuclear reactor. The bacterium realigns its radiation-shattered pieces of DNA. Then enzymes bring in new nucleotides and assemble the pieces.

The discovery of DNA repair systems began with observations in the late 1940s that when fungi were exposed to ultraviolet (UV) radiation, those cultures later placed nearest a window grew best. The researchers who noted these effects were not investigating DNA repair, but were using UV light in other experiments. Therefore, DNA repair was inadvertently discovered before the structure of DNA was. The DNA-damaging effect of UV radiation, and the ability of light to correct it, was soon observed in a variety of organisms. (UV radiation has a shorter wavelength than visible light. They are both types of electromagnetic radiation.)

Types of DNA Repair

Exposure to radiation is a fact of life. Since its beginning, the Earth has been periodically bathed in UV radiation. Volcanoes, comets, meteorites, and supernovas all depleted ozone in the atmosphere, which allowed ultraviolet wavelengths of light to reach organisms. The shorter wavelengths—UVA—are not dangerous, but the longer UVB wavelengths damage DNA by forming an extra covalent bond between adjacent (same-strand) pyrimidines, particularly thymines (**figure 12.13**). The linked thymines are called thymine dimers. Their extra bonds kink the double helix sufficiently to disrupt replication and permit insertion of a noncomplementary base. For example, an A might be inserted opposite a G or C, instead of opposite a T. Thymine dimers also disrupt transcription.

Early in the evolution of life, organisms that could survive UV damage had an advantage. Enzymes enabled them to do this, and because enzymes are gene-encoded, DNA repair came to persist.

In many modern species, three types of DNA repair peruse the genetic material for mismatched base pairs. In the first type, enzymes called photolyases absorb energy from visible light and use it to detect and bind to pyrimidine dimers, then break the extra bonds. This type of repair, called photo-reactivation, enables UV-damaged fungi to recover from exposure to sunlight. Humans do not have this type of DNA repair.

In the early 1960s, researchers discovered a second type of DNA self-mending, called **excision repair**, in mutant *E. coli* that were unable to repair UV-induced DNA damage. Enzymes cut the bond between the DNA sugar and base and snip out—or excise—the pyrimidine dimer and surrounding

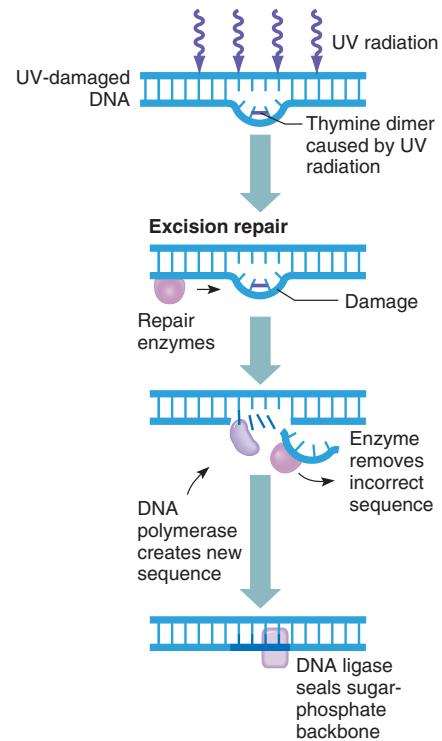


Figure 12.13 Excision repair. Human DNA damaged by UV light is repaired by excision repair, which removes and replaces the pyrimidine dimer and a few surrounding bases.

bases (see figure 12.13). Then, a DNA polymerase fills in the correct nucleotides, using the exposed template as a guide. DNA polymerase also detects and corrects mismatched bases in newly replicated DNA.

Humans have two types of excision repair. **Nucleotide excision repair** replaces up to 30 nucleotides and removes errors that result from several types of insults, including exposure to chemical carcinogens, UVB in sunlight, and oxidative damage. Thirty different proteins carry out nucleotide excision repair, functioning together as a structure called a repairosome.

The second type of excision repair, **base excision repair**, replaces one to five nucleotides at a time, but specifically corrects errors that result from oxidative damage. Oxygen free radicals are highly reactive forms of oxygen that arise during chemical reactions such as those of metabolism and transcription. Free radicals damage DNA. Genes that are very actively transcribed face greater oxidative damage from free radicals; base excision repair targets this type of damage.

A third mechanism of DNA sequence correction is **mismatch repair**. Enzymes “proofread” newly replicated DNA for small loops that emerge from the double helix. The enzymes excise the mismatched base so that it can be replaced (**figure 12.14**). These loops emerge from where the two strands do not precisely align, but instead slip and misalign.

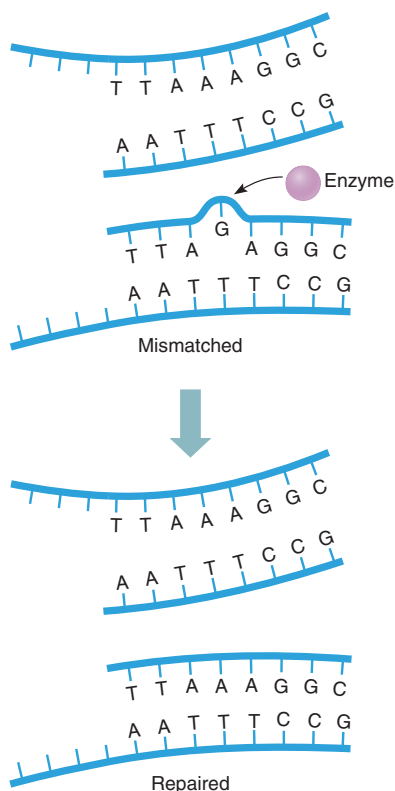


Figure 12.14 Mismatch repair. In this form of DNA repair, enzymes detect loops and bulges in newly replicated DNA that indicate mispairing. The enzymes correct the error. Highly repeated sequences are more prone to this type of error.

This occurs where very short DNA sequences repeat. These sequences, called microsatellites, are scattered throughout the genome. Like minisatellites, microsatellite lengths can vary from person to person, but within an individual, they are usually the same length. Excision and mismatch repair differ in the cause of the error—UV-induced pyrimidine dimers versus replication errors—and in the types of enzymes involved.

The three forms of DNA repair in human cells relieve the strain on thymine dimers or replace incorrectly inserted bases. Another form of repair can heal a broken sugar-phosphate backbone in both strands, which can result from exposure to ionizing radiation or oxidative damage. This type of insult breaks a chromosome, which can cause cancer. At least two types of multiprotein complexes reseal the sugar-phosphate backbone, either by rejoining the broken ends or recombining with DNA on the unaffected homolog.

In yet another type of DNA repair called damage tolerance, a “wrong” DNA base is left in place, but replication and transcription proceed. “Sloppy” DNA polymerases, with looser adherence to the base-pairing rules, read past the error, randomly inserting any other base. It is a little like retaining a misspelled word in a sentence—usually the meaning remains clear.

Figure 12.15 summarizes DNA repair mechanisms.

DNA Repair Disorders

The ability to repair DNA is crucial to health. If both copies of a repair gene are mutant, a disorder can result. Heterozygotes who have one mutant repair gene may be more sensitive to damage from environmental toxins.

A well-studied DNA damage response gene encodes a protein called p53. It controls whether DNA is repaired and the cell salvaged, or the cell dies by apoptosis (see figure 18.15). Signal transduction activates p53 protein, stabilizing and aggregating it into complexes of four proteins. These quartets bind DNA at four palindromic repeats that indicate genes that slow the cell cycle. The cycle must slow for repair to take place. If the damage is too severe, the p53 protein quartets instead increase the rate of transcription of genes that promote apoptosis.

In DNA repair disorders, chromosome breakage caused by factors such as radiation cannot be repaired. Mutations in repair genes therefore greatly increase susceptibility to certain types of cancer following exposure to ionizing radiation or chemicals that affect cell division. These conditions develop because errors in the DNA sequence accumulate and are perpetuated to a much greater extent than they are in people with functioning repair systems. We conclude this chapter with a closer look at repair disorders.

Trichothiodystrophy (MIM 601675)

At least five genes can cause trichothiodystrophy. At its worst, this condition causes dwarfism, mental retardation, and failure to develop, in addition to brittle hair and scaly skin, both with low sulfur content. Although the child may appear to be normal for a year or two, growth soon slows dramatically, signs of premature aging begin, and life ends early. Hearing and vision may fail. Interestingly, the condition does not increase the risk of cancer. Symptoms reflect accumulating oxidative damage. Individuals have faulty nucleotide excision repair, base excision repair, or both.

Inherited Colon Cancer

Hereditary nonpolyposis colon cancer (HNPCC) (MIM 120435 and others, also known as Lynch syndrome) was linked to a DNA repair defect when researchers discovered different-length microsatellites within an individual. Because mismatch repair normally keeps a person’s microsatellites all the same length, people with this type of colon cancer might have a breakdown in this form of DNA repair. HNPCC is common, affecting 1 in 200 people. Mutations in any of at least 7 genes can cause this form of colon cancer.

HNPCC accounts for 3 percent of newly diagnosed cases of colorectal cancer. Genetic testing for this condition in all people newly diagnosed with colon cancer is advised because if they have a mutation, their relatives can be tested. If healthy relatives test positive, frequent colonoscopies can detect disease early, at a more treatable stage. Penetrance of HNPCC is about 45 percent by age 70—considered a high cancer risk.

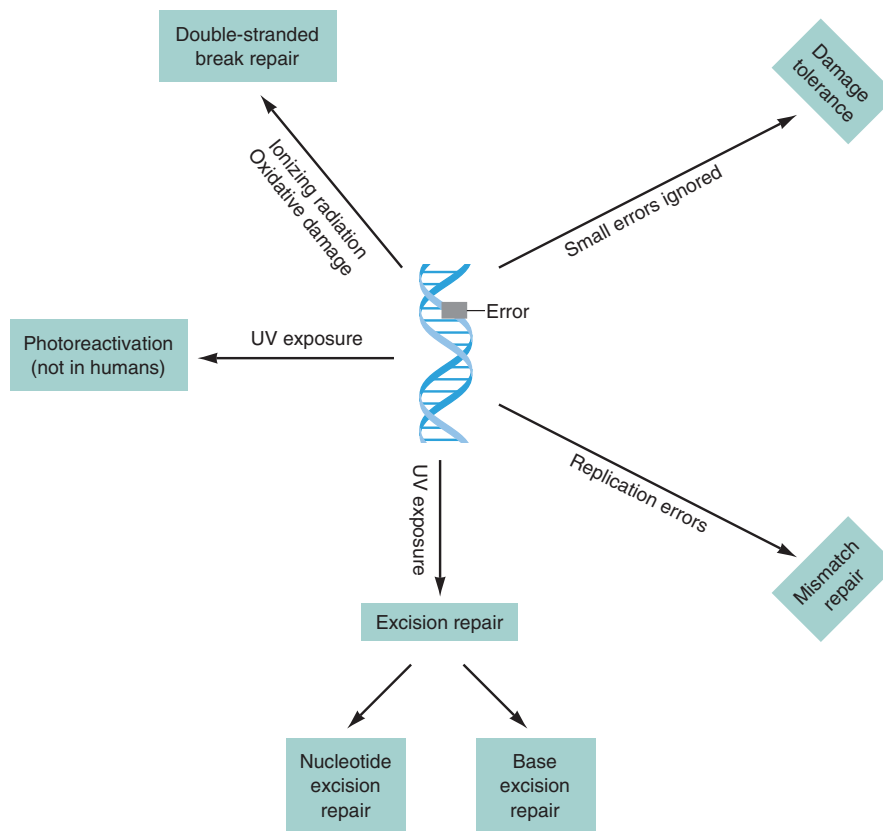


Figure 12.15 DNA repair mechanisms.

Xeroderma Pigmentosum (XP) (MIM 278700)

A child with XP must stay indoors in artificial light, because even the briefest exposure to sunlight causes painful blisters. Failing to cover up and use sunblock can result in skin cancer (figure 12.16). More than half of all children with XP develop the cancer before they reach their teens. People with XP have a 1,000-fold increased risk of developing skin cancer compared to others, and a 10-fold increase in the risk of developing internal tumors.

XP is autosomal recessive, and results from mutations in any of seven genes. It can reflect malfunction of nucleotide excision repair or deficient “sloppy” DNA polymerase, both of which allow thymine dimers to stay and block replication.

One of the genes that causes XP, when mutant, also causes trichothiodystrophy and another disorder, Cockayne syndrome. The different symptoms arise from the different ways that mutations disrupt the encoded protein, which is a helicase that helps unwind replicating DNA.

Only about 250 people in the world are known to have XP. A family living in upstate New York runs a special summer camp for children with XP, where they turn night into day. Activities take place at night, or in special areas where the windows are covered and light comes from low-ultraviolet incandescent lightbulbs.

Ataxia Telangiectasis (AT) (MIM 208900)

This multisymptom disorder is the result of a defect in a kinase that functions as a cell cycle checkpoint (see figure 2.15). In AT, cells proceed through the cell cycle without pausing just after replication to inspect the new DNA and to repair any mispaired bases. Some cells die through apoptosis if the damage is too great to repair. Because of the malfunctioning cell cycle, individuals who have this autosomal recessive disorder have 50 times the risk of developing cancer, particularly of the blood. About 40 percent of individuals with ataxia telangiectasis have cancer by age 30. Additional symptoms include poor balance and coordination (ataxia), red marks on the face (telangiectasia), delayed sexual maturation, and high risk of contracting lung infections and developing diabetes mellitus. These symptoms probably arise from disruption of other functions of the kinase.

AT is rare, but heterozygotes are not. They make up from 0.5 to 1.4 percent of various populations. Carriers may have mild radiation sensitivity, which causes a two- to sixfold increase in cancer risk over the general population. People who know they are AT carriers should question suggested dental or medical X rays, because for them even low exposure to radiation may cause cancer.

DNA’s changeability, so vital for evolution of a species, comes at the cost of occasional harm to individuals. Each of us harbors a few mutations, and several polymorphisms, although most are hidden in the recessive state. Individuals whose mutations cause illness or deformity can face hardships, both medical and due to discrimination. Perhaps we can learn from the ancient Egyptians, who honored people who were genetically different.

The dry air of ancient Egypt and the meticulous recording of daily life in art and burial places reveal that people with short stature, particularly those with autosomal dominant achondroplasia (a form of dwarfism), were accepted, important, and even revered as gods. Ancient Egyptian “little people” were jewelers, animal keepers, entertainers, and personal attendants, often to royalty. High-ranking people with dwarfism were given special burial places near the pyramids. Wrote Chahira Kozma, a professor of pediatrics at Georgetown University who studies how the ancient Egyptians regarded unusual people, “Dwarfs were accepted in ancient Egypt; their recorded daily activities suggest assimilation into daily life, and their disorder was



Figure 12.16 A DNA repair disorder. The marks on this child's face result from sun exposure. He is highly sensitive because he has inherited xeroderma pigmentosum (XP), an impairment of excision repair. The large lesion on his chin is a skin cancer.

not shown as a physical handicap. Wisdom writings and moral teachings in ancient Egypt commanded respect for dwarfs and other individuals with disabilities.”

Today, photographer Rick Guidotti continues the tradition of honoring genetic diversity in his moving images of people with inherited disease. He contributed figure 4.17*b*. Visit his website at positiveexposure.org.

Key Concepts

1. Many genes encode enzymes that locate and correct errors in replicating DNA, in the DNA damage response.
2. A common cause of noncomplementary base insertion is a UV-induced pyrimidine dimer.
3. Photoreactivation or excision repair can unkink pyrimidine dimers.
4. Mismatch repair corrects noncomplementary base pairs that are inserted into newly replicated DNA.
5. Repair seals broken sugar-phosphate backbones.
6. DNA damage tolerance allows replication to proceed past a mismatched base.
7. Abnormal repair genes cause disorders usually associated with chromosome breaks and predisposition to cancer.

Summary

12.1 The Nature of Mutations

1. A **mutation** is a change in a gene's nucleotide base sequence that affects less than 1 percent of a population and can cause a **mutant** phenotype. A polymorphism is more common and may not alter the phenotype.
2. A **germline mutation** originates in meiosis and affects all cells of an individual. A **somatic mutation** originates in mitosis and affects a subset of cells.
3. A mutation disrupts the function or abundance of a protein or introduces a new function. Loss-of-function mutations are usually recessive, and altered or gain-of-function mutations are dominant.

12.2 Two Mutations

4. Mutations in the beta globin and collagen genes cause a variety of disorders.
5. Whether different mutations in a gene cause the same or distinct illnesses varies; nomenclature is inconsistent.

12.3 Causes of Mutation

6. A spontaneous mutation arises due to chemical phenomena or to an error in DNA replication. Spontaneous mutation rate is characteristic of a gene and is more likely in repeats.
7. **Mutagens** are chemicals or radiation that delete, substitute, or add bases. An organism may be exposed to a mutagen intentionally, accidentally, or naturally.

12.4 Types of Mutations

8. A **point mutation** alters a single DNA base. It may be a **transition** (purine to purine or pyrimidine to pyrimidine) or a **transversion** (purine to pyrimidine or vice versa). A **missense mutation** substitutes one amino acid for another, while a **nonsense mutation** substitutes a “stop” codon for a codon that specifies an amino acid, shortening the protein product. Point mutations in splice sites can add or delete amino acids.
9. Adding or deleting genetic material may upset the reading frame or otherwise alter protein function.
10. A pseudogene results when a duplicate of a gene mutates. It may disrupt chromosome pairing, causing mutation.
11. Transposons may disrupt the functions of genes they jump into.
12. Expanding triplet repeat mutations add stretches of the same amino acid to a protein. They expand because they attract each other, which affects replication.
13. **Copy number variants** are DNA sequences that are repeated a different number of times in different individuals. They may have no effect on phenotype or may directly or indirectly cause disease.

12.5 The Importance of Position

14. Several types of mutations can affect a gene.
15. Mutations in the globin genes may affect the ability of the blood to transport oxygen, or they may have no effect.

16. Susceptibility to prion disorders requires two mutations that affect different parts of the protein that interact as the amino acid chain folds.
 17. Synonymous codons limit the effects of mutation. Changes in the second codon position often substitute a similarly shaped amino acid.
 18. **Conditional mutations** are expressed only in response to certain environmental triggers.
 19. Sending the most recently replicated DNA into cells headed for differentiation, while sending older strands into stem cells, protects against mutation.
- ### 12.6 DNA Repair
20. DNA polymerase proofreads DNA, but repair enzymes correct errors in other ways.
 21. Photoreactivation repair uses light energy to split pyrimidine dimers.
 22. In **excision repair**, pyrimidine dimers are removed and the area filled in correctly. **Nucleotide excision repair** replaces up to 30 nucleotides from various sources of mutation. **Base excision repair** fixes up to five bases that paired incorrectly due to oxidative damage.
 23. **Mismatch repair** proofreads newly replicated DNA for loops that indicate noncomplementary base pairing.
 24. DNA repair also fixes the sugar-phosphate backbone. Damage tolerance enables replication to continue beyond a mismatch.
 25. Mutations in repair genes break chromosomes and increase cancer risk.

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Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

1. How do a “silent mutation” and a polymorphism that is a “harmless variant” differ?
2. How does a point mutation differ from a SNP, and how are they similar?
3. Distinguish between a germline and a somatic mutation. Which is likely to be more severe? Which can be transmitted to offspring?
4. Why is the collagen molecule especially likely to be altered by mutation?
5. What criteria should be used to determine whether mutations in a gene are likely to cause different disorders or differing degrees of the same disorder?
6. How can DNA spontaneously mutate?
7. Compare the effects of alpha, beta, and gamma radiation on the human body.
8. What is the physical basis of a mutational hot spot?
9. List two types of mutations that can alter the reading frame.
10. List four ways that DNA can mutate without affecting the phenotype.
11. Cite two ways a jumping gene can disrupt gene function.
12. What is a molecular explanation for the worsening of an inherited illness over generations?
13. Compare and contrast how short repeats within a gene, long triplet repeats within a gene, and repeated genes can cause disease.
14. How does a copy number variant differ from a missense mutation?
15. Why can a mutation that retains an intron’s sequence and a triplet repeat mutation have a similar effect on a gene’s encoded protein?
16. Why is there not “a” human genome, but many?
17. Cite three ways in which the genetic code protects against the effects of mutation.
18. What is a conditional mutation?
19. How do excision and mismatch repair differ?
20. In trichothiodystrophy, brittle hair and scaly skin arise in some patients only during periods of fever that persist long enough for hair, nail, and skin changes to become noticeable. What type of mutation causes this disorder?
21. Explain how semiconservative DNA replication makes it possible for stem cells to receive the DNA least likely to bear mutations.
22. Consult the genetic code (figure 10.12).
 - a. Describe a point mutation that would not affect a protein’s primary structure.
 - b. Look up amino acid structures and identify a point mutation that could greatly alter a protein.

Applied Questions

- Consider the following sequence of part of an mRNA molecule:

AUGUUGUCAAAAGCAUGGCGGCCA

 Introduce the following changes to the sequence, and indicate the effect, if any, on the encoded amino acid sequence:
 - a missense mutation
 - a nonsense mutation
 - a frameshift mutation
 - a silent mutation
 - a transversion
 - a transition
 - a tandem duplication
 - a deletion
- Retinitis pigmentosa causes night blindness and loss of peripheral vision before age 20. A form of X-linked retinitis pigmentosa (MIM 300455) is caused by a frameshift mutation that deletes 199 amino acids. How can a simple mutation have such a great effect?
- A mutation that changes a C to a T causes a form of Ehlers-Danlos syndrome, forming a “stop” codon and shortened procollagen. Consult the genetic code and suggest one way that this can happen.
- Part of the mRNA sequence of an exon of a gene that encodes a blood protein is:

AUGACUCAUCGCGUGUAGUUUACGA

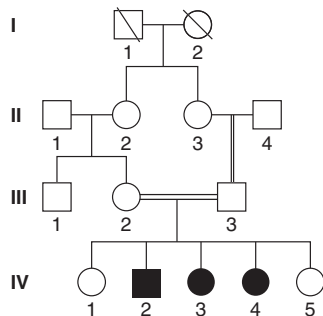
 Consult the genetic code to answer the following questions:
 - What is the sequence of amino acids that this mRNA encodes?
 - What is the sequence if a point mutation changes the tenth base from a C to an A?
 - What is the effect of a point mutation that changes the fifteenth base from a U to an A?
 - How does the encoded amino acid sequence change if a C is inserted between the fourth and fifth bases?
 - Which would be more devastating to the encoded amino acid sequence, insertion of three bases in a row, or insertion of two bases in a row?
- Susceptibility to developing prion diseases entails a mutation from aspartic acid (*asp*) to asparagine (*asn*). Which nucleotide base changes make this happen?
- Two teenage boys meet at a clinic to treat muscular dystrophy. The boy who is more severely affected has a two-base insertion at the start of his dystrophin gene. The other boy has the same two-base insertion but also has a third base inserted a few bases away. Explain why the second boy’s illness is milder.
- Two missense mutations in the gene that encodes an enzyme called superoxide dismutase cause amyotrophic lateral sclerosis (ALS, or Lou Gehrig’s disease). This disorder causes loss of neurological function over a 5-year period. One mutation alters the amino acid asparagine (*asn*) to lysine (*lys*). The other changes an isoleucine (*ile*) to a threonine (*thr*). List the codons involved and describe how single-base mutations alter the amino acids they specify.
- In one family, Tay-Sachs disease stems from a four-base insertion, which changes an amino-acid-encoding codon into a “stop” codon. What type of mutation is this?
- Epidermolytic hyperkeratosis (MIM 607602) is an autosomal dominant condition that produces scaly skin. It can be caused by a missense mutation that substitutes a histidine (*his*) amino acid for an arginine (*arg*). Write the mRNA codons that could account for this change.
- A point mutation in the gene *CRYBA4* (MIM 123631) replaces a phenylalanine codon with a serine codon. The gene encodes β -crystallin protein, discussed in section 5.1. In a healthy eye, the proteins align, sculpting the lens. When the gene is mutant, the proteins cannot align and vision is impaired. Consult the genetic code and suggest two ways that a point mutation can account for this amino acid substitution.
- Aniridia (MIM 106200) is an autosomal dominant eye condition in which the iris is absent. In one family, an 11-base insertion in the gene causes a very short protein to form. What kind of mutation must the insertion cause?
- A biotechnology company has encapsulated DNA repair enzymes in fatty bubbles called liposomes. Why would this be a valuable addition to a tanning lotion?

Web Activities

- Children with Hutchinson-Gilford progeria syndrome (figure 3.22) age extremely rapidly. In 18 of 20 children, a single base change in the *lamin A* gene alters a C to a T, but this mutation removes 50 amino acids from the encoded protein. In all 20 children, the parents do not have the mutation.
 - Is the mutation in the 18 children *de novo* or induced? What is the evidence for this distinction?
 - How can a change in a single base remove 50 amino acids?
 - Using MIM, list and describe six other disorders caused by mutation in the *lamin A* gene.
- Go to http://www.lpaonline.org/resources_faq.html and read the Little People of America’s Position Statement on Genetic Discoveries in Dwarfism. Cite an application of genetic testing for achondroplasia that could be construed as beneficial, and one that could be thought of as harmful.
- Select an image at positiveexposure.org and describe the symptoms of the photographed disorder.

Case Studies and Research Results

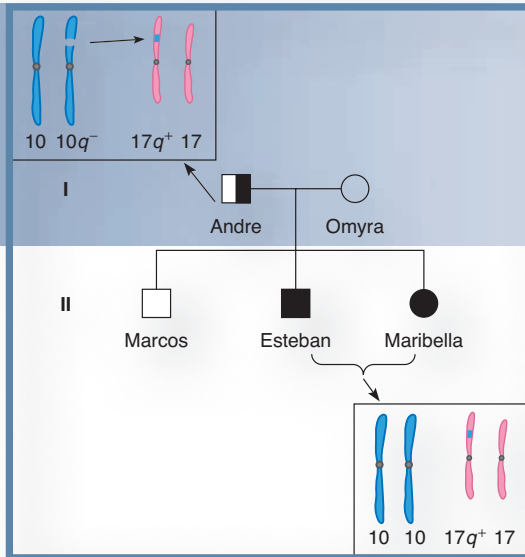
16. Latika and Keyshauna meet at a clinic for college students who have cystic fibrosis. Latika's mutation results in exon skipping. Keyshauna's mutation is a nonsense mutation. Which young woman probably has more severe symptoms? Cite a reason for your answer.
17. Marshall and Angela have skin cancer resulting from xeroderma pigmentosum. They meet at an event for teenagers with cancer. However, their mutations affect different genes. They decide to marry but not to have children because they believe that each child would have a 25 percent chance of inheriting XP because it is autosomal recessive. Are they correct? Why or why not?
18. For several disorders, copy number variants represent a small fraction of patients. For any of the following such disorders, describe the copy number variant mutation or any other causative mutation by consulting MIM:
- DiGeorge syndrome
 - Williams-Beuren syndrome
 - spinal muscular atrophy (any type)
 - schizophrenia
 - Smith-Magenis syndrome
 - Prader-Willi syndrome
 - Angelman syndrome
 - nephronophthisis
 - β -crystallin cataracts
 - psoriasis
19. Two girls and a boy in a Pakistani family have a form of deafness caused by a mutation in the gene that encodes a protein called tricellulin (MIM 610153). The normal protein attaches epithelial (lining) cells in groups of three in the inner ear in a way that establishes compartments whose fluid content differences are crucial to hearing. Below is a pedigree for the family.



- What is the mode of inheritance for this form of deafness, and how do you know this?
 - This form of deafness is rare worldwide, but more common among Pakistani families, many of whose pedigrees have double horizontal lines like the parents in the third generation of this pedigree. What does the double line mean, and how does it account for the increased prevalence of this form of deafness in the population?
 - The affected children have the following partial sequence for the tricellulin gene: C T G C A A T G T. Unaffected family members have the corresponding sequence: C T G C A G T G T. What are the amino acid differences encoded in these sequences?
20. Presenilin 1 is one of the genes that, when mutant, causes familial Alzheimer disease. The gene is expressed in the heart, and certain mutations cause a condition called dilated cardiomyopathy that leads to heart failure. In the Esposito family, all of the relatives who have or had heart failure have the following partial sequence for the presenilin 1 gene: G A T G A T G G C G G G. Family members with healthy hearts have the sequence G A T G G T G G C G G G. How do the encoded amino acid sequences differ between the healthy and sick family members for this part of the gene?
21. At the Center for Applied Genomics at the Children's Hospital of Philadelphia, 100,000 young children are having their genomes scanned for copy number variants. Researchers are comparing the resulting CNV profiles with multifactorial disorders that the children have, such as asthma, obesity, cancers, and diabetes, to determine whether copy number variation is important to health. The idea is that the children are so young that environmental influences are minimized, compared, for example, to people who have asthma but have smoked or lived with pollution. Explain how this project might be helpful and how it might be harmful.

Forensics Focus

22. Late one night, a man broke into Colleen's apartment and raped her. He wore a mask and it was dark, so she couldn't see his face, but she did manage to yank out some of his long, greasy hair. The forensic investigator, after examining the hair, asked Colleen if her boyfriend had been in the bed or with her earlier in the evening, because the hairs were of two genotypes for one of the repeated sequences that was analyzed.
- What is another explanation for the finding of two genotypes?
 - How can investigators distinguish between the two explanations for finding cells of two genotypes in the evidence?



Too much genetic material. A piece of the short arm of Andre's chromosome 10 (10q⁻) has moved to one chromosome 17. He is healthy. However, Esteban and Maribella have each inherited the copy of the chromosome 17 with the extra material (17q⁺) as well as two normal chromosome 10s. The extra chromosome 10 DNA caused their symptoms.

CHAPTER

13

Chromosomes

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13.5 Uniparental Disomy—A Double Dose from One Parent**A LATE DIAGNOSIS**

Esteban was 17 years old when he learned that he had an unusual chromosome. The discovery explained a lot.

As a baby and toddler, Esteban had been much slower to walk and talk than his brother Marcos. When he started school, problems emerged: He was learning disabled and had difficulty interacting with others. Esteban had to repeat the third grade so that he could learn to read. He was also very tall and thin, causing teachers to think he was older than he really was and therefore to expect more of him. Still, Esteban was able to learn in regular classrooms with weekly visits to a resource room, and he made a few friends.

Esteban's mother, Omyra, became pregnant when Esteban was in the tenth grade. She hadn't had a fetal chromosome check when she was pregnant with Esteban, but this time she did. The test showed one chromosome 17 with a bit of another chromosome stuck into it. A geneticist at the medical center where Omyra's doctor practiced requested DNA samples from the family. She found that the fetus had inherited the unusual chromosome from the father, Andre, but the chromosome charts from father and fetus differed. Andre's cells had two unusual chromosomes but a normal amount of genetic material. A bit of chromosome 10 had inserted into chromosome 17, and the fetus received only the unusual chromosome 17 from Andre. Esteban's chromosomes had the same abnormality as his future sister's. His symptoms may have stemmed from the extra DNA. His little sister Maribella was slow, too, but is today a very happy preschooler.

13.1 Portrait of a Chromosome

Mutations range from single-base changes to entire extra sets of chromosomes. A mutation is considered a chromosomal aberration if it is large enough to see with a light microscope using stains and/or fluorescent probes to highlight missing, extra, or moved genetic material.

In general, excess genetic material has milder effects on health than a deficit. Still, most large-scale chromosomal abnormalities present in all cells disrupt or halt prenatal development. As a result, only 0.65 percent of all newborns have chromosomal abnormalities that produce symptoms. An additional 0.20 percent have chromosomal rearrangements in which chromosome parts have been flipped or swapped, but they do not produce symptoms unless they disrupt genes that are crucial to health.

Cytogenetics is the subdiscipline within genetics that links chromosome variations to specific traits, including illnesses. This chapter explores several chromosome-level abnormalities and their effects on health. Actual cases are used to describe some of them.

Required Parts: Telomeres and Centromeres

A chromosome consists primarily of DNA and proteins with a small amount of RNA, and is duplicated and transmitted—via mitosis or meiosis—to the next cell generation. Chromosomes have long been described and distinguished by size and shape, using stains and dyes to contrast dark **heterochromatin** with the lighter **euchromatin** (figure 13.1). Heterochromatin consists mostly of highly repetitive DNA sequences, whereas euchromatin has more protein-encoding sequences.

A chromosome must include structures that enable it to replicate and remain intact—everything else is essentially

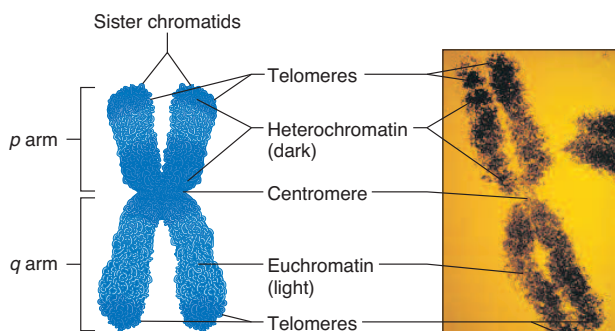


Figure 13.1 Portrait of a chromosome. Tightly wound, highly repetitive heterochromatin forms the centromere (the largest constriction) and the telomeres (the tips) of chromosomes. Elsewhere, lighter-staining euchromatin includes many protein-encoding genes. The centromere divides this chromosome into a short arm (*p*) and a long arm (*q*). This chromosome is in the replicated form.

informational cargo (protein-encoding genes and their controls). The essential parts of a chromosome are:

- telomeres
- origin of replication sites, where replication forks begin to form
- the centromere

Recall from figure 2.18 that **telomeres** are chromosome tips. In humans, each telomere is many repeats of the sequence TTAGGG. In most cell types, telomeres shorten with each mitotic cell division.

The **centromere** is the largest constriction of a chromosome and it is where spindle fibers attach when the cell divides. A chromosome without a centromere is no longer a chromosome. It vanishes from the cell as soon as division begins because there is no way to attach to the spindle.

Centromeres, like chromosomes, are made up mostly of DNA and protein. Many of the hundreds of thousands of DNA bases that form the centromere are repeats of a specific 171-base DNA sequence. The size and number of repeats are similar in many species, although the sequence differs. This suggests that these repeats have a structural role in maintaining chromosomes rather than an informational role. Certain centromere-associated proteins are synthesized only when mitosis is imminent, forming a structure called a kinetochore that contacts the spindle fibers, enabling the cell to divide.

Centromeres are replicated toward the end of S phase. A protein that may control their duplication is called centromere protein A, or CENP-A. Molecules of CENP-A stay with centromeres as chromosomes are replicated, covering about half a million DNA base pairs. When the replicated (sister) chromatids separate at anaphase, each member of the pair retains some CENP-A. The protein therefore passes to the next cell generation, but it is *not* DNA. This is another example of an epigenetic change.

Centromeres lie within vast stretches of heterochromatin. The arms of the chromosome extend outward from the centromere. Gradually, the DNA includes more protein-encoding sequences as distance from the centromere increases. Gene density varies greatly among chromosomes. Chromosome 21 is a gene “desert,” harboring a million-base stretch with no protein-encoding genes at all. Chromosome 22, in contrast, is a gene “jungle.” These two tiniest chromosomes are remarkably similar in size, but chromosome 22 contains 545 genes to chromosome 21’s 225!

The chromosome parts that lie between protein-rich areas and the telomeres are termed subtelomeres (figure 13.2). These areas extend from 8,000 to 300,000 bases inward toward the centromere from the telomeres. Subtelomeres include some protein-encoding genes and therefore bridge the gene-rich regions and the telomere repeats. The transition is gradual. Areas of 50 to 250 bases, right next to the telomeres, consist of 6-base repeats, many of them very similar to the TTAGGG of the telomeres. Then, moving inward from the 6-base zone are many shorter repeats, each present in a few copies. Their

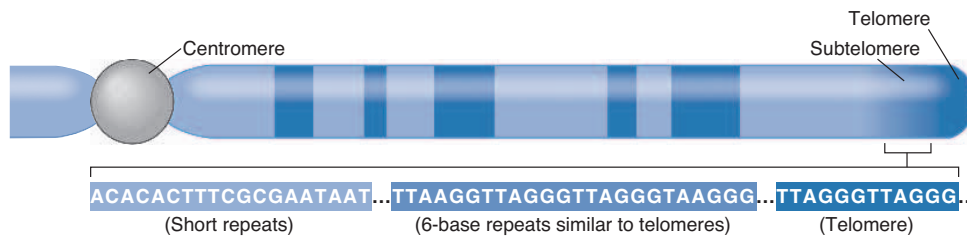


Figure 13.2 Subtelomeres. The repetitive sequence of a telomere gradually diversifies toward the centromere. The centromere is depicted as a buttonlike structure to more easily distinguish it, but it is composed of DNA like the rest of the chromosome.

function isn't known. Finally the sequence diversifies and protein-encoding genes appear.

At least 500 protein-encoding genes lie in the total subtelomere regions. About half are members of multigene families (groups of genes of very similar sequence next to each other) that include pseudogenes. These multigene families may reflect recent evolution: Apes and chimps have only one or two genes for many of the large gene families in humans. Such gene organization is one explanation for why our genome sequence is so very similar to that of our primate cousins—but we are clearly different animals. Our genomes differ more in gene copy number and chromosomal organization than in DNA base sequence.

Karyotypes Chart Chromosomes

Even in this age of genomics, the standard chromosome chart, or **karyotype**, remains a major clinical tool. A karyotype displays chromosomes in pairs by size and by physical landmarks that appear during mitotic metaphase, when DNA coils tightly. **Figure 13.3** shows a karyotype with an extra chromosome.

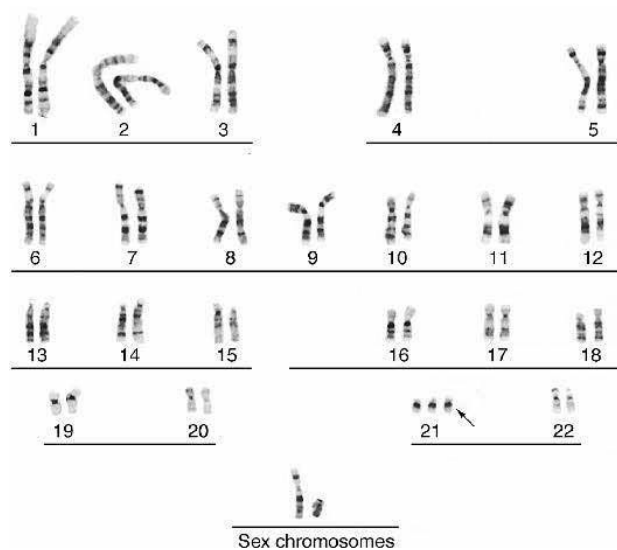


Figure 13.3 A karyotype displays chromosome pairs in size order. Note the extra chromosome 21 that causes trisomy 21 Down syndrome.

The 24 human chromosome types are numbered from largest to smallest—1 to 22. The other two chromosomes are the X and the Y. Early attempts to size-order chromosomes resulted in generalized groupings because many of the chromosomes are of similar size. Use of dyes and stains made it easier to distinguish chromosomes because they form patterns of bands.

Centromere position is one physical feature of chromosomes. A chromosome is **metacentric** if the centromere divides it into two arms of approximately equal length. It is **submetacentric** if the centromere establishes one long arm and one short arm, and **acrocentric** if it pinches off only a small amount of material toward one end (**figure 13.4**). Some species have telocentric chromosomes that have only one arm, but humans do not. The long arm of a chromosome is designated *q*, and the short arm *p* (*p* stands for “petite”).

Five human chromosomes (13, 14, 15, 21, and 22) have bloblike ends, called satellites, that extend from a thinner, stalklike bridge from the rest of the chromosome. The stalk regions do not bind stains well. The stalks carry many copies of genes encoding ribosomal RNA and ribosomal proteins. These areas coalesce to form the nucleolus, a structure in the nucleus where ribosomal building blocks are produced and assembled (see figure 2.3).

Karyotypes are useful at several levels. When a baby is born with the distinctive facial features of Down syndrome, a karyotype confirms the clinical diagnosis. Within families,

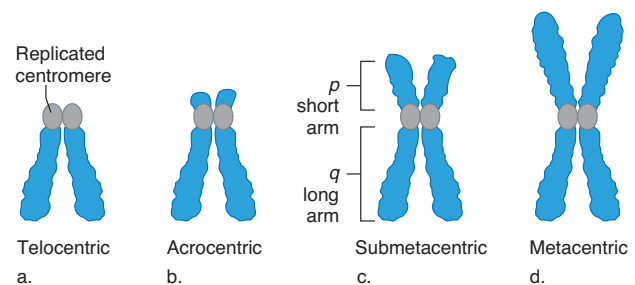


Figure 13.4 Centromere position distinguishes chromosomes. (a) A telocentric chromosome has the centromere toward one end although telomere DNA sequences are still at the tip. Humans do not have any telocentric chromosomes. (b) An acrocentric chromosome has the centromere near an end. (c) A submetacentric chromosome's centromere creates a long arm (*q*) and a short arm (*p*). (d) A metacentric chromosome's centromere establishes equal-sized arms.

karyotypes are used to identify relatives with a particular chromosomal aberration that can affect health. In one family, several adults died from a rare form of kidney cancer. Karyotypes revealed that the affected individuals all had an exchange, called a **translocation**, between chromosomes 3 and 8. When karyotypes showed that two healthy young family members had the translocation, physicians examined and monitored their kidneys. Cancer was found very early and successfully treated.

Karyotypes of individuals from different populations can reveal the effects of environmental toxins, if abnormalities appear only in a group exposed to a particular contaminant. Because chemicals and radiation that can cause cancer and birth defects often break chromosomes into fragments or rings, detecting this genetic damage can alert physicians to the possibility that certain cancers may appear in the population.

Karyotypes compared among species can clarify evolutionary relationships. The more recent the divergence of two species from a common ancestor, the more closely related we presume they are, and the more alike their chromosome banding patterns should be. Our closest relative, according to karyotypes, is the pygmy chimpanzee (bonobo). The human karyotype is also remarkably similar to that of the domestic cat, and somewhat less similar to those of mice, pigs, and cows. Among mammals, it is least like the karyotype of the aardvark, indicating that this is a primitive placental mammal.

Key Concepts

1. A chromosome minimally includes telomeres, origins of replication, and centromeres.
2. A centromere consists of DNA repeats and associated proteins, some of which bind spindle fibers. Centromere protein A enables the centromere to replicate.
3. Subtelomeres contain telomeric repeats and protein-encoding multigene families.
4. Chromosomes differ by size, centromere location, satellites, and staining. Karyotypes are size-order chromosome charts.

13.2 Visualizing Chromosomes

Extra or missing chromosomes are detected by counting a number other than 46. Identifying chromosome rearrangements, such as an inverted sequence or an exchange of parts between two chromosomes, requires a way to distinguish among the chromosomes. A combination of stains and DNA probes applied to chromosomes allows this. A **DNA probe** is a labeled piece of DNA that binds to its complementary base sequence on a particular chromosome.

Obtaining Cells for Chromosome Study

Any cell other than a mature red blood cell (which lacks a nucleus) can be used to examine chromosomes, but some cells are easier to obtain and culture than others. Skinlike cells

collected from the inside of the cheek are the easiest to obtain for a chromosome test; white blood cells are used too. A person might require a chromosome test if he or she has a family history of a chromosomal abnormality or seeks medical help because of infertility.

Chromosome tests are commonly performed on cells from fetuses. Couples who receive a prenatal diagnosis of a chromosome abnormality can arrange for treatment of the newborn, if possible; learn more about the condition and contact support groups and plan care; or terminate the pregnancy. These choices are best made after a genetic counselor or physician provides information on the medical condition and treatment options.

Chromosomes of a fetus are checked in several ways. **Amniocentesis** and **chorionic villus sampling** have been available for many years. They sample fetal cells from the amniotic fluid and chorionic villi, respectively, and detect large-scale chromosomal abnormalities. A newer technique called **chromosome microarray analysis** can be paired with the older techniques to detect copy number variants, which include extremely small sections of missing or extra DNA. Chromosome microarray analysis probes and displays specific sequences, detecting many disorders that other techniques miss.

Amniocentesis

The first fetal karyotype was constructed in 1966 using amniocentesis. In this procedure, a doctor removes a small sample of fetal cells and fluids from the uterus with a needle passed through the woman's abdominal wall (**figure 13.5a**). The cells are cultured for a week to 10 days, and typically 20 cells are karyotyped. The sampled amniotic fluid may also be examined for deficient, excess, or abnormal biochemicals that could indicate an inborn error of metabolism. Tests for specific single-gene disorders are based on family history and may be done on cells in the amniotic fluid sample as well. Ultrasound is used to follow the needle's movement and to visualize fetal parts, such as the profile in **figure 13.6**.

Amniocentesis can detect approximately 1,000 of the more than 5,000 known chromosomal and biochemical problems. The most common chromosomal abnormality detected is one extra chromosome, called a **trisomy**. Amniocentesis is usually performed between 14 and 16 weeks gestation, when the fetus isn't yet very large but amniotic fluid is plentiful. Amniocentesis can be carried out anytime after this point.

Doctors recommend amniocentesis if the risk that the fetus has a detectable condition exceeds the risk that the procedure will cause a miscarriage. Until recently, this risk cutoff was thought to be about age 35 in the woman, when the risk to the fetus of a detectable chromosome problem about equals the risk of amniocentesis causing pregnancy loss—1 in 350. While it is still true that the risk of a chromosomal problem rises steeply after maternal age 35, amniocentesis has become much safer in the 30 or so years since the statistics were obtained that have been used for most risk estimates (**figure 13.7**). In 2007 a large study found the risk of amniocentesis causing miscarriage to be about 1 in 1,600, leading some physicians and organizations

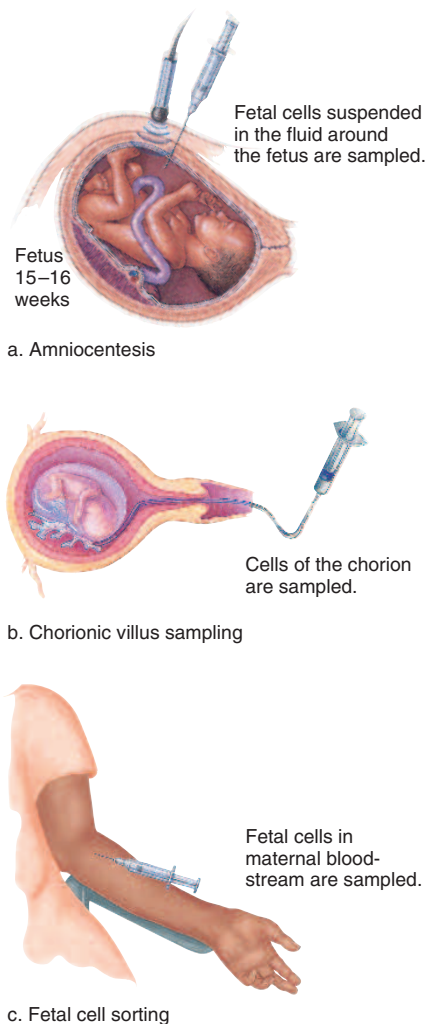


Figure 13.5 Three ways to check a fetus's chromosomes.

(a) Amniocentesis draws out amniotic fluid. Fetal cells shed into the fluid are collected and their chromosomes examined. (b) Chorionic villus sampling removes cells that would otherwise develop into the placenta. Since these cells descended from the fertilized ovum, they should have the same chromosomal constitution as the fetus. (c) Researchers can detect fetal cells, DNA, or mRNA in a sample of blood from a pregnant woman.

to offer amniocentesis to younger women too. The procedure is also warranted if a couple has had several spontaneous abortions or children with birth defects or a known chromosome abnormality, irrespective of maternal age.

Another reason to seek amniocentesis is if screening tests on a pregnant woman indicate elevated risk for a trisomy (extra chromosome) of the fetus. These “multiple maternal serum marker” tests, discussed in *Bioethics: Choices for the Future* on page 245, are offered to all pregnant women. Cut-off levels for the results based on population statistics are used to identify fetuses at elevated risk, and the women are then offered



Figure 13.6 A sonogram is an image obtained with ultrasound. In an ultrasound exam, sound waves bounced off the embryo or fetus are converted into a three-dimensional appearing image. “4D ultrasound” provides a video of an embryo or fetus. (The fourth dimension is time.)

the more invasive amniocentesis and chorionic villus sampling, which yield a definitive diagnosis. Screening tests consider maternal age, ultrasound findings, and levels of certain proteins in the woman’s blood at certain times in the pregnancy.

Chorionic Villus Sampling

During the 10th through 12th week of pregnancy, chorionic villus sampling (CVS) obtains cells from the chorionic villi, which are finger-like structures that develop into the placenta (figure 13.5b). A karyotype is prepared directly from the collected cells, rather than first culturing them, as in amniocentesis. Results are ready in days.

Because chorionic villus cells descend from the fertilized ovum, their chromosomes should be identical to those of the embryo and fetus. Occasionally, a chromosomal aberration

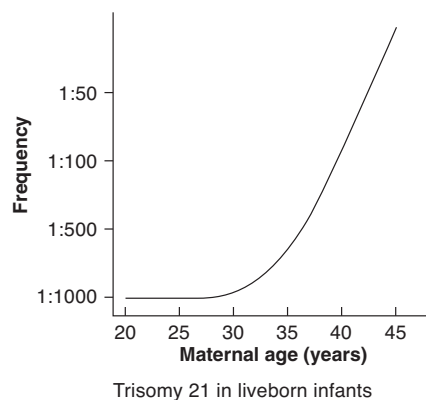


Figure 13.7 The risk of conceiving an offspring with trisomy 21 rises dramatically with maternal age.

Bioethics: Choices for the Future

The Denmark Study: Screening for Down Syndrome

Prenatal tests for trisomy 21 Down syndrome are of two general types. *Screening tests* identify fetuses that are at increased risk of having trisomy 21. These tests consider an ultrasound finding (excess fluid at the back of the neck) and abnormal levels of certain proteins in the pregnant woman's blood (multiple maternal serum markers) and maternal age. (**table 1**). Screening tests that include serum markers are routinely offered to pregnant women of any age, and much more accurately predict risk than maternal age alone.

If screening tests find a fetus to be at elevated risk for trisomy 21, more invasive *diagnostic tests* are offered—chorionic villus sampling or amniocentesis. Both are highly accurate, but introduce a small risk of miscarriage. In contrast to the screening tests, the diagnostic tests actually visualize the extra chromosome.

In 2004, the government of Denmark issued new guidelines offering prenatal screening for trisomy 21 to all pregnant women. Earlier guidelines offered diagnostic tests only to women over age 35, based on assessment by age alone and using old statistics. Informed consent was required—that is, all women who took the screening tests knew the risks and benefits.

Researchers have tracked number of trisomy 21 births and use of diagnostic testing (CVS and amniocentesis) in Denmark

from 2000 until 2006 to assess impact of the new guidelines. The number of infants born with trisomy 21 has been halved, the number diagnosed before birth increased by 30 percent, and the number of invasive prenatal diagnostic tests done each year has decreased by 50 percent.

Health care is nationalized in Denmark, and prenatal screening for trisomy 21 is available to all. In addition, all of the nation's nineteen departments of obstetrics and gynecology use the same software for their record keeping, so that it is easy to maintain national databases of test results and clinical outcomes. Studies are ongoing to assess the societal and bioethical repercussions from implementing the new guidelines.

Questions for Discussion

1. What is a medical benefit of the guidelines in Denmark?
2. What do you think potential patients should be told during the informed consent process?
3. Mark Leach, chairman of the Down Syndrome Affiliates in Action group, said that testing has "outpaced society's understanding of what life with Down syndrome is like." How would you feel about the program if you had a child with Down syndrome?
4. How can the Danish government prevent women from feeling pressured to have prenatal testing for trisomy 21?
5. Explain how the number of infants born with Down syndrome declined more than half, but additional prenatal diagnosis increased only by about a third.
6. Down syndrome patients are living longer and richer lives than they have in the past. How can this fact be reconciled with a population-wide screening program such as the one in Denmark?
7. What should the role of the father be in deciding whether a woman should have prenatal screening or diagnosis for trisomy 21?

Table 1

Commonly Used Maternal Serum Markers

Marker	Elevated Risk
Alpha fetoprotein (AFP)	< normal
Human chorionic gonadotropin (hCG)	> normal
Estriol	< normal
Inhibin A	> normal
Pregnancy-associated plasma protein A	< normal

occurs only in a cell of the embryo, or only in a chorionic villus cell. This results in chromosomal mosaicism—the karyotype of a villus cell differs from that of an embryo cell. Chromosomal mosaicism has great clinical consequences. If CVS indicates an abnormality in villus cells that is not also in the fetus, then a couple may elect to terminate the pregnancy when the fetus is actually chromosomally normal. In the opposite situation, the results of the CVS may be normal, but the fetus has abnormal chromosomes.

CVS is slightly less accurate than amniocentesis, and in about 1 in 1,000 to 3,000 procedures, it halts development of the feet and/or hands, a condition termed transverse limb defects. Also, CVS does not sample amniotic fluid, so tests for

inborn errors of metabolism are not possible. The advantage of CVS is earlier results, but the disadvantage is a greater risk of spontaneous abortion. However, CVS has become much safer in recent years.

Fetal Cells, DNA, and RNA

Detecting fetal cells or nucleic acids in the pregnant woman's bloodstream is safer than amniocentesis and CVS, but is still experimental in the United States (figure 13.5c). The technique traces its roots to 1957, when a pregnant woman died when cells from a very early embryo lodged in a major blood vessel in her lung, blocking blood flow. The fetal cells were detectable because

they were from a male, and contained the telltale Y chromosome. This meant that fetal cells could enter a woman's circulation.

By studying the blood of other pregnant women, researchers found that fetal cells enter the maternal circulation in up to 70 percent of pregnancies. Cells from female embryos, however, cannot easily be distinguished from the cells of the pregnant woman on the basis of sex chromosome analysis. But fetal cells from either sex can be distinguished from maternal cells using a device called a fluorescence-activated cell sorter. It separates fetal cells from maternal blood by identifying surface characteristics that differ from those on the woman's cells. The fetal cells are then karyotyped and specific gene tests performed on fetal DNA.

Free fetal DNA is also in a pregnant woman's bloodstream, but is difficult to detect because it is so rare. A new technique detects fetal mRNA in the woman's bloodstream. If fetal mRNA transcribed from a specific sequence that differs between the parents is not present in equal amounts, then presence of an extra copy of the sequence in the fetus is inferred. If the sequence is part of chromosome 21, for example, the technique can diagnose the extra chromosome of Down syndrome.

Preparing Cells for Chromosome Observation

Cytogeneticists have tried to describe and display human chromosomes since the late nineteenth century (**figure 13.8**). Then, the prevailing view held that humans had an XO sex determination system, with females having an extra chromosome (XX). Estimates of the human chromosome number ranged from 30 to 80. In 1923, Theophilus Painter published sketches of human chromosomes from three patients at a Texas state mental hospital. The patients had been castrated in an attempt to control their abusive behavior, and Painter was able to examine the removed tissue. He could not at first tell whether the cells had 46 or 48 chromosomes, but finally decided that he saw 48. Painter later showed that both sexes have the same chromosome number.

The difficulty in distinguishing between 46 or 48 chromosomes was physical—it is challenging to prepare a cell in which chromosomes do not overlap. To easily count the chromosomes, scientists had to find a way to capture them when they are most condensed—during cell division—and also spread them apart. Since the 1950s, cytogeneticists have used colchicine, an extract of the crocus plant, to arrest cells during division.

Swelling, Squashing, and Untangling

How to untangle the spaghetti-like mass of chromosomes was solved by accident in 1951. A technician mistakenly washed white blood cells being prepared for chromosome analysis in a salt solution that was less concentrated than the interiors of the cells. Water rushed into the cells, swelling them and separating the chromosomes. Then cell biologists Albert Levan and Joe-Hin Tjio found that drawing cell-rich fluid into a pipette and dropping it onto a microscope slide prepared with stain burst the cells and freed the mass of chromosomes. Adding a glass coverslip spread the chromosomes enough that they could be counted. Researchers finally could see that the number of chromosomes in a diploid human cell is 46, and that the number in gametes is 23.

Karyotypes were once constructed using a microscope to locate a cell where the chromosomes were not touching, photographing the cell, developing a print, cutting out the individual chromosomes, and arranging them into a size-ordered chart. Today, a computer scans ruptured cells in a drop of stain and selects one in which the chromosomes are the most visible and well-spread. Then image analysis software recognizes the band patterns of each stained chromosome pair, sorts the structures into a size-ordered chart, and prints the karyotype. If the software recognizes an abnormal band pattern, a database pulls out identical or similar karyotypes from records of other patients.

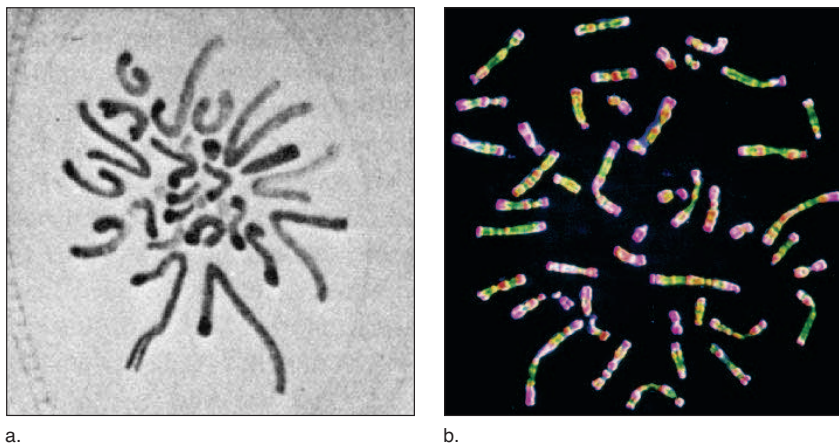


Figure 13.8 Viewing chromosomes, then and now. (a) The earliest drawings of chromosomes, by German biologist Walter Flemming, date from 1882. His depiction captures the random distribution of chromosomes as they splash down on a slide. (b) A micrograph of actual stained human chromosomes.

Staining

In the earliest karyotypes, dyes were used to stain chromosomes a uniform color. Chromosomes were grouped into size classes, designated A through G, in decreasing size order. In 1959, scientists described the first chromosomal abnormalities—Down syndrome (an extra chromosome 21), Turner syndrome (also called XO syndrome, a female with only one X chromosome), and Klinefelter syndrome (also called XXY syndrome, a male with an extra X chromosome). Visualizing and distinguishing the sex chromosomes revealed the causes of these conditions, discussed later in the chapter.

The first chromosome stains could highlight large deletions and duplications, but usually researchers only vaguely understood the nature of a chromosomal syndrome. In 1967, a mentally retarded child with material missing from chromosome 4 would have been diagnosed as having a “B-group chromosome” disorder.

Describing smaller chromosomal aberrations required better ways to distinguish chromosomes. In the 1970s, Swedish scientists developed stains that create banding patterns unique to each chromosome. These stains are specific for AT-rich or GC-rich stretches of DNA, or for heterochromatin, which is dark-staining. A band represents at least 5 to 10 million DNA bases.

The ability to detect missing, extra, inverted, or misplaced bands allowed researchers to link many more syndromes with specific chromosome aberrations. Then researchers found that synchronizing the cell cycle of cultured cells revealed more bands per chromosome. Another technique, **fluorescence *in situ* hybridization** (FISH), introduced the ability to highlight individual genes.

FISHing

FISH is more precise and targeted than conventional chromosome staining because it uses DNA probes that are complementary to specific DNA sequences. The probes are attached to molecules that fluoresce when illuminated, producing a flash of color precisely where the probe binds to a chromosome in a patient’s sample.

FISH can “paint” entire karyotypes by probing each chromosome with several different fluorescent molecules. A computer integrates the images and creates a unique false color for each chromosome. Many laboratories that perform amniocentesis or CVS use FISH probes for chromosomes 13, 18, 21, and the X and Y to quickly identify the most common problems. In **figure 13.9**, FISH reveals the extra chromosome 21 in cells from a fetus with trisomy 21 Down syndrome.

A new type of prenatal chromosome analysis amplifies certain repeated sequences on chromosomes 13, 18, 21, X, and Y. The technique distinguishes paternally derived from maternally derived repeats on each homolog for these five chromosomes. An abnormal ratio of maternal to paternal repeats indicates a numerical problem, such as two copies of one parent’s chromosome 21. Combined with the one chromosome 21 from the

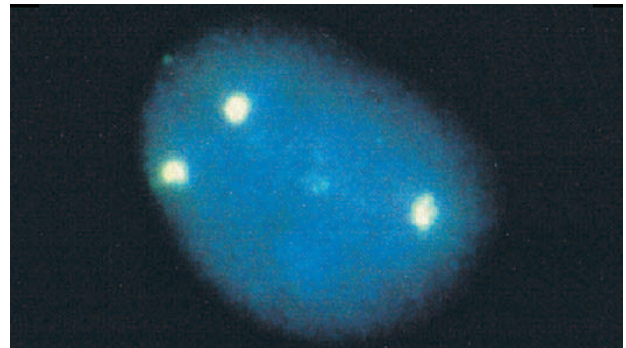


Figure 13.9 FISHing for genes and chromosomes. FISH shows three fluorescent dots that correspond to three copies of chromosome 21. Each dot represents a specific DNA sequence to which the fluorescently-labeled probe complementary base pairs.

other parent, this situation would produce a fertilized ovum with three copies of chromosome 21, which causes Down syndrome.

Chromosomal Shorthand

Geneticists abbreviate the pertinent information in a karyotype by listing chromosome number, then sex chromosome constitution, then abnormal autosomes. Symbols describe the type of aberration, such as a deletion or translocation; numbers correspond to specific bands. A normal male is 46,XY; a normal female is 46,XX. Geneticists use band notation to describe gene locations. For example, the gene encoding the β -globin subunit of hemoglobin is located at 11p15.5. **Table 13.1** gives some examples of chromosomal shorthand.

Chromosome information is displayed in a graphical representation called an ideogram. (**figure 13.10**). The chromosome is divided into arms and numbered regions and

Table 13.1 Chromosomal Shorthand

Abbreviation	What It Means
46,XY	Normal male
46,XX	Normal female
45,X	Turner syndrome (female)
47,XXY	Klinefelter syndrome (male)
47,XYY	Jacobs syndrome (male)
46,XY, del (7q)	A male missing part of the long arm of chromosome 7
47,XX, + 21	A female with trisomy 21 Down syndrome
46,XY, t(7;9)(p21.1; q34.1)	A male with a translocation between the short arm of chromosome 7 at band 21.1 and the long arm of chromosome 9 at band 34.1
48, XYY	A male with an extra X and an extra Y

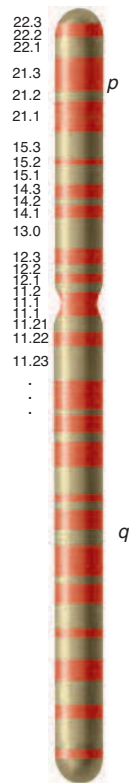


Figure 13.10 Ideogram. An ideogram is a schematic chromosome map. It indicates chromosome arm (*p* or *q*) and major regions delineated by banding patterns. This figure is repeated in the context of more specific ways to depict chromosomes in figure 22.1.

subregions. Specific gene loci are sometimes listed on the right side. Ideograms are becoming so crowded with notations indicating specific genes that they may soon become obsolete.

Key Concepts

1. Karyotypes display chromosomes in size order.
2. Chromosomes can be visualized in any cell that has a nucleus and can be cultured.
3. Fetal karyotypes are made from cells obtained by amniocentesis, CVS, or fetal cell sorting from maternal blood. Maternal serum marker patterns and ultrasound scans can reveal increased risk of an abnormal chromosome number.
4. Cytogeneticists obtain cells; display, stain, and probe chromosomes with fluorescent molecules; and then arrange them in a karyotype.
5. Chromosomal shorthand summarizes the number of chromosomes, sex chromosome constitution, and type of aberration. Ideograms display features of individual chromosomes.

13.3 Abnormal Chromosome Number

A human karyotype is abnormal if the number of chromosomes in a somatic cell is not 46, or if individual chromosomes have extra, missing, or rearranged genetic material.

Abnormal chromosomes account for at least 50 percent of spontaneous abortions, yet only 0.65 percent of newborns have abnormal chromosomes. Therefore, most embryos and fetuses with abnormal chromosomes stop developing before birth. **Table 13.2** summarizes the types of chromosome abnormalities in the order in which they are discussed.

Improved technology has made it possible to detect extremely small amounts of extra or missing genetic material and, as a result, more people are being diagnosed with chromosomal abnormalities. Today, many families whose members have the same chromosome abnormalities are finding each other through organizations on the Internet, communicating by e-mail or meeting in person, and sharing experiences. **Table 13.3** lists some of these organizations. Often families with the same chromosomal disorder then form organizations, such as the “International 22q11.2 Deletion Syndrome Foundation,” a group of families with members missing DNA on the short arm of chromosome 22.

Polyploidy

The most drastic upset in chromosome number is an entire extra set. A cell with extra sets of chromosomes is **polyploid**. An individual whose cells have three copies of each chromosome is a triploid (designated 3N, for three sets of chromosomes). Two-thirds of all triploids result from fertilization of an oocyte by two sperm. The other cases arise from formation of a diploid gamete, such as when a normal haploid sperm fertilizes

Table 13.2 Chromosome Abnormalities

Type of Abnormality	Definition
Polyploidy	Extra chromosome sets
Aneuploidy	An extra or missing chromosome
Monosomy	One chromosome absent
Trisomy	One chromosome extra
Deletion	Part of a chromosome missing
Duplication	Part of a chromosome present twice
Translocation	Two chromosomes join long arms or exchange parts
Inversion	Segment of chromosome reversed
Isochromosome	A chromosome with identical arms
Ring chromosome	A chromosome that forms a ring due to deletions in telomeres, which cause ends to adhere

Table 13.3 Organizations for Families with Chromosome Abnormalities

Hope for Trisomy 13 + 18	www.hopefortrisomy13and18.org/
Rainbows Down Under	http://members.optushome.com.au/karens/
Support Organization for Trisomy 18, 13 and Related Disorders (SOFT)	http://www.trisomy.org/
Tracking Rare Incidence Syndromes (TRIS)	http://web.coehs.siu.edu/Grants/TTRIS/
UNIQUE—the Rare Chromosome Disorder Support Group	http://www.rarechromo.org/html/home.asp

a diploid oocyte. Triploids account for 17 percent of spontaneous abortions (**figure 13.11**). Very rarely, an infant survives as long as a few days, with defects in nearly all organs. However, certain human cells may be polyploid. The liver, for example, has some tetraploid (4N) and even octaploid (8N) cells.

Polyploids are very common among flowering plants, including roses, cotton, barley, and wheat, and in some insects. Fish farmers raise triploid salmon, which cannot breed.

Aneuploidy

Cells missing a single chromosome or having an extra one are **aneuploid**, which means “not good set.” Rarely, aneuploids can have more than one missing or extra chromosome, indicating defective meiosis in a parent. A normal chromosome number is **euploid**, which means “good set.”

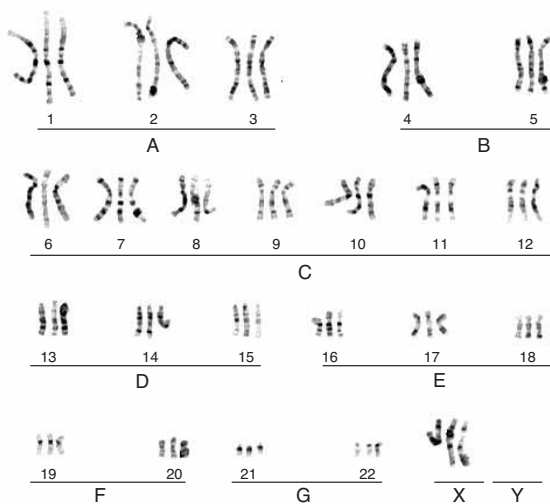


Figure 13.11 Polyploids in humans are nearly always lethal. Individuals with three copies of each chromosome (triploids) in every cell account for 17 percent of all spontaneous abortions and 3 percent of stillbirths and newborn deaths.

Most autosomal aneuploids (with a missing or extra non-sex chromosome) are spontaneously aborted. Those that survive have specific syndromes, with symptoms depending upon which chromosomes are missing or extra. Mental retardation is common in aneuploidy because development of the brain is so complex and of such long duration that nearly any chromosome-scale disruption affects genes whose protein products affect the brain. Sex chromosome aneuploidy usually produces milder symptoms.

Most children born with the wrong number of chromosomes have an extra chromosome (a trisomy) rather than a missing one (a monosomy). Most monosomies are so severe that an affected embryo ceases developing. Trisomies and monosomies are named for the chromosomes involved, and in the past the associated syndromes were named for the discoverers. Today, cytogenetic terminology is used because it is more precise. For example, Down syndrome can result from a trisomy or a translocation. The distinction is important in genetic counseling. Translocation Down syndrome, although accounting for only 4 percent of cases, has a much higher recurrence risk within a family than the trisomy form, a point we return to later in the chapter.

The meiotic error that causes aneuploidy is called **nondisjunction**. Recall that in normal meiosis, homologs separate and each of the resulting gametes receives only one member of each chromosome pair. In nondisjunction, a chromosome pair fails to separate at anaphase of either the first or second meiotic division. This produces a sperm or oocyte that has two copies of a particular chromosome, or none, rather than the normal one copy (**figure 13.12**). When such a gamete fuses with its partner at fertilization, the zygote has either 45 or 47 chromosomes, instead of the normal 46. Different trisomies tend to be caused by nondisjunction in the male or female, at meiosis I or II.

A cell can have a missing or extra chromosome in 49 ways—an extra or missing copy of each of the 22 autosomes, plus the five abnormal types of sex chromosome combinations—Y, X, XXX, XXY, and XYY. (Sometimes individuals have four or even five sex chromosomes.) However, only nine types of aneuploids are recognized in newborns. Others are seen in spontaneous abortions or fertilized ova intended for *in vitro* fertilization.

Most of the 50 percent of spontaneous abortions that result from extra or missing chromosomes are 45,X individuals (missing an X chromosome), triploids, or trisomy 16. About 9 percent of spontaneous abortions are trisomy 13, 18, or 21. More than 95 percent of newborns with abnormal chromosome numbers have an extra 13, 18, or 21, or an extra or missing X or Y chromosome. These conditions are all rare at birth—together they affect only 0.1 percent of all children. But nondisjunction occurs in 5 percent of recognized pregnancies.

Types of chromosome abnormalities seem to differ between the sexes. Abnormal oocytes mostly have extra or missing chromosomes, whereas abnormal sperm more often have structural variants, such as inversions or translocations, discussed later in the chapter.

Aneuploidy and polyploidy also arise during mitosis, producing groups of somatic cells with the extra or missing chromosome. An individual with two chromosomally distinct

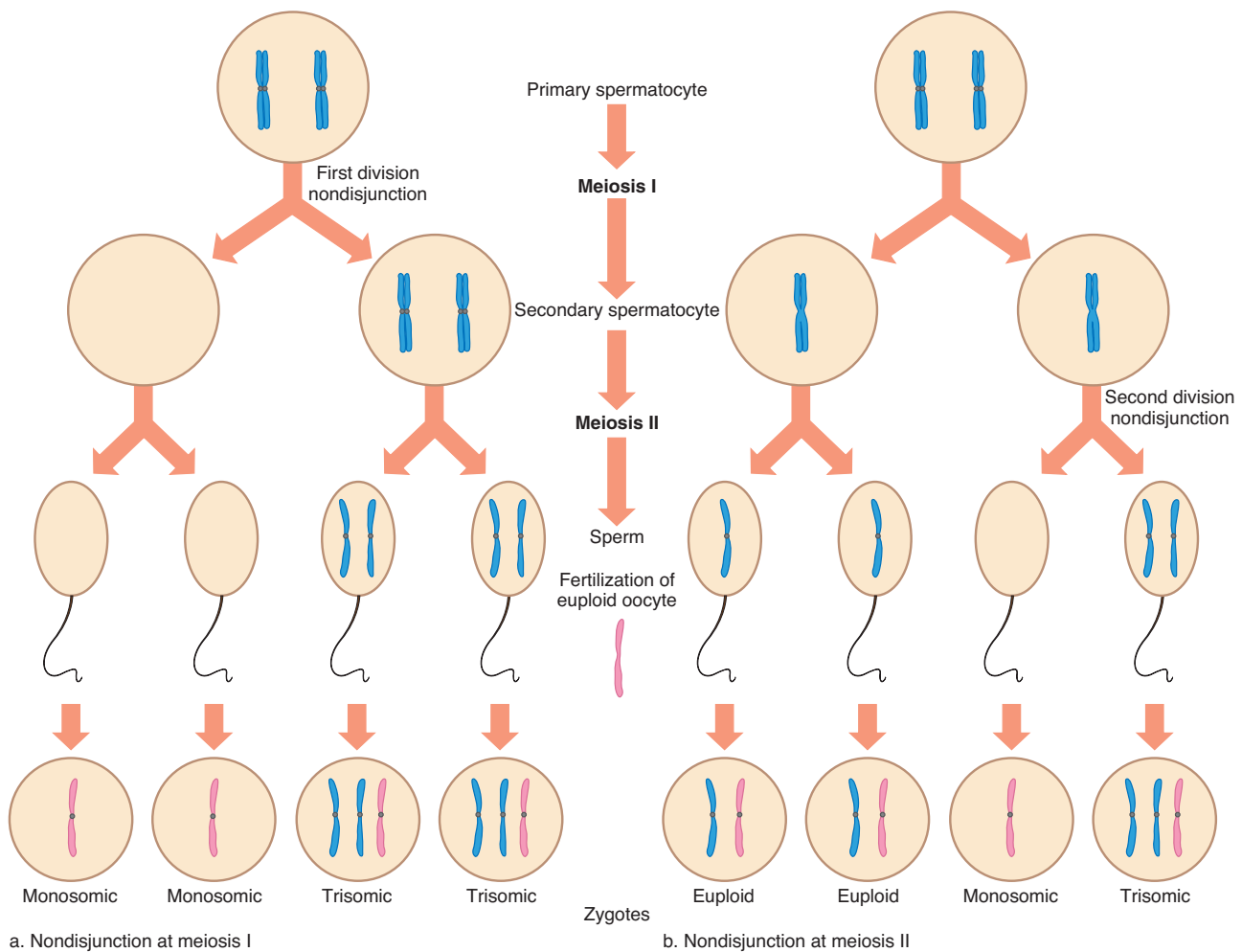


Figure 13.12 Extra and missing chromosomes—aneuploidy. Unequal division of chromosome pairs can occur at either the first or second meiotic division. **(a)** A single pair of chromosomes is unevenly partitioned into the two cells arising from meiosis I in a male. The result: two sperm cells have two copies of the chromosome, and two sperm cells have no copies. When a sperm cell with two copies of the chromosome fertilizes a normal oocyte, the zygote is trisomic; when a sperm cell lacking the chromosome fertilizes a normal oocyte, the zygote is monosomic. **(b)** This nondisjunction occurs at meiosis II. Because the two products of the first division are unaffected, two of the mature sperm are normal and two are aneuploid. Oocytes can undergo nondisjunction as well, leading to zygotes with extra or missing chromosomes when normal sperm cells fertilize them.

cell populations is a mosaic. If only a few cells are altered, health may not be affected. However, a mitotic abnormality that occurs early in development, so that many cells descend from the unusual one, can affect health. A chromosomal mosaic for a trisomy may have a mild version of the associated condition. This is usually the case for the 1 to 2 percent of people with Down syndrome who are mosaic. The phenotype depends upon which cells have the extra chromosome. Unfortunately, prenatal testing cannot reveal which cells are affected.

Autosomal Aneuploids

Most autosomal aneuploids cease developing long before birth. Following are cases and descriptions of the most common autosomal aneuploids among liveborns. The most frequently seen

extra autosomes in newborns are chromosomes 21, 18, and 13 because these chromosomes carry far fewer protein-encoding genes than the other autosomes, compared to their total amount of DNA. Therefore, extra copies of these chromosomes are tolerated well enough for some fetuses with them to survive to be born (**table 13.4**).

Trisomy 21—David's Story

When David G. was born in 1994, doctors told his 19-year-old mother, Toni, to put him into an institution. "They said he wouldn't walk, talk, or do anything. Today, I want to bring him back and say look, he walks and talks and runs track and is graduating high school," says Toni.

Like other teens, David has held part-time jobs, gone to dances, and uses a computer. But he is unlike most other teens

Table 13.4 Comparing and Contrasting Trisomies 13, 18, and 21

Type of Trisomy	Incidence at Birth	Percent of Conceptions That Survive 1 Year After Birth
13 (Patau)	1/12,500–1/21,700	<5%
18 (Edward)	1/6,000–1/10,000	<5%
21 (Down)	1/800–1/826	85%

in that his cells have an extra chromosome 21, which limits his intellectual abilities. “Maybe he’s not book smart, but when you look around at what he can do, he’s smart,” Toni says. His speech is difficult to understand, and he has facial features characteristic of Down syndrome, but he has a winning personality and close friends.

Sometimes David gets into unusual situations because he takes things literally. He once dialed 911 when he stubbed his toe, because he’d been told to do just that when he was hurt. Overall he’s done so well that he plans to move into a group home and find a job after high school.

Still, Toni fears that he will meet few people like himself as he gets older. She read a report that 90 percent of couples told their fetus has Down syndrome end the pregnancy.

The most common autosomal aneuploid among liveborns is trisomy 21. The extra folds in the eyelids, called epicanthal folds, and flat face prompted Sir John Langdon Haydon Down to term the condition *mongoloid* when he described it in 1866. As the medical superintendent of a facility for the profoundly mentally retarded, Down noted that about 10 percent of his patients resembled people of Mongolian heritage. The resemblance is superficial. People of all ethnic groups are affected.

A person with Down syndrome is usually short and has straight, sparse hair and a tongue protruding through thick lips. The hands have an abnormal pattern of creases, the joints are loose, and poor reflexes and muscle tone give a “floppy” appearance. Developmental milestones (such as sitting, standing, and walking) come slowly, and toilet training may take several years. Intelligence varies greatly. Parents of a child with Down syndrome can help their child reach maximal potential by providing a stimulating environment. Some people with Down syndrome can attend college and hold jobs (figure 13.13).

Many people with Down syndrome have physical problems, including heart and kidney defects and hearing and vision loss. A suppressed immune system can make influenza deadly. Digestive system blockages are common and may require surgical correction. A child with Down syndrome is 15 times more likely to develop leukemia than a child who does not have the syndrome, but this is still only a 1 percent risk. Many of the medical problems associated with Down syndrome are treatable, so that average life expectancy is now in the fifties. In 1910, life expectancy was only 9 years.

Some people with Down syndrome who pass age 40 develop the black fibers and tangles of amyloid beta protein in their brains characteristic of Alzheimer disease, although they



Figure 13.13 Trisomy 21. Many years ago, people with Down syndrome could be institutionalized. Today, thanks to tremendous strides in both medical care and special education, some people with the condition can graduate college.

usually do not become severely demented (see Reading 5.1). The chance of a person with trisomy 21 developing Alzheimer disease is 25 percent, compared to 6 percent for the general population. A gene on chromosome 21 causes one inherited form of Alzheimer disease. Perhaps the extra copy of the gene in trisomy 21 has a similar effect to a mutation in the gene that causes Alzheimer disease, such as causing amyloid beta buildup. However, Alzheimer disease is not inevitable for people with Down syndrome. A medical journal chronicled the case of “Mr. C,” a 70-year-old with Down syndrome who does not have any signs of dementia at all.

Before the human genome sequence became available, researchers studied people who have a third copy of only part of chromosome 21 to identify specific genes that could cause symptoms. Table 13.5 lists some genes known to contribute to trisomy 21 Down syndrome symptoms.

The likelihood of giving birth to a child with trisomy 21 Down syndrome increases dramatically with the age of the mother (see figure 13.7). However, 80 percent of children with trisomy 21 are born to women under age 35, because younger women are more likely to become pregnant and less likely to have amniocentesis. About 90 percent of trisomy 21 conceptions are due to nondisjunction during meiosis I in the female. The 10 percent of cases due to the male result from nondisjunction during meiosis I or II. The chance that trisomy 21 will recur in a family, based on empirical data (how often it actually does recur in families), is 1 percent.

The age factor in trisomy 21 Down syndrome and other trisomies may reflect the fact that the older a woman is, the longer her oocytes have been arrested on the brink of completing meiosis. This is a time period of 15 to 45 years, when oocytes may have been exposed to toxins, viruses, and radiation. A second explanation for the maternal age effect is that females have a pool of immature aneuploid oocytes resulting from spindle abnormalities that cause nondisjunction. As a woman ages, selectively releasing normal oocytes each month, the abnormal ones remain, much as black jelly beans accumulate as people

Table 13.5 Genes Associated with Trisomy 21 Down Syndrome

Gene Product	MIM	Signs and Symptoms (Phenotype)
Amyloid precursor protein (APP)	104760	Protein deposits in brain
Chromatin assembly factor I (CAF1A)	601245	Impaired DNA synthesis
Collagen type VI (COL6A1)	120220	Heart defects
Crystallin (CRYA1)	123580	Cataracts
Cystathione beta synthase (CBS)	236200	Impaired metabolism and DNA repair
Interferon receptor 1 (IFNAR)	107450	Impaired immunity
Kinase 1 (DYRK1A)	600855	Mental retardation
Oncoprotein ETS2 (ETS2)	164740	Skeletal abnormalities, cancer
Phosphoribosylglycinamide formyltransferase (GART)	138440	Impaired DNA synthesis and repair
Superoxide dismutase (SOD1)	147450	Premature aging

preferentially eat the colored ones. Yet a third possible explanation for the maternal age effect is that trisomies result from gametes in which a homolog pair do not extensively cross over during meiosis I. Such chromosomes tend to migrate to the same pole, packaging an extra chromosome into a gamete.

The association between maternal age and Down syndrome has been recognized for a long time, because affected individuals were often the youngest children in large families. Before the chromosome connection was made in 1959, the syndrome was thought to be caused by syphilis, tuberculosis, thyroid malfunction, alcoholism, or emotional trauma, or even “maternal reproductive exhaustion.” The increased risk

of Down syndrome correlates to maternal age, not to the number of children in the family.

Trisomy 18—Anthony’s Story

When an ultrasound scan early in pregnancy revealed a small fetus with low-set ears, a small jaw, a pocket of fluid in the brain, and a peculiarly clenched fist, the parents-to-be, Elisa and Brendan, were advised to have amniocentesis to view the fetus’s chromosomes. The signs on the scan suggested an extra chromosome 18. Amniocentesis confirmed what the ultrasound suggested.

Although Elisa and Brendan were stunned and upset to learn what lay ahead, they continued the pregnancy. The fetus remained small, as Elisa swelled hugely with three times the normal volume of amniotic fluid. Further ultrasound scans revealed that only one of the baby’s kidneys worked, the heart had holes between the chambers, and part of the intestine lay outside the stomach in a sac, so tube feed-

ing would be necessary. The child would be severely developmentally delayed and mentally retarded. Anthony was delivered at 36 weeks after his heart rate became erratic during a routine prenatal visit. He lived only 22 days.

Trisomies 18 and 13 were described in the same research report in 1960 (figure 13.14). Trisomy 18 is also called Edward syndrome and trisomy 13 is also known as Patau syndrome. Most affected individuals do not survive to be born.

Children who have trisomy 18 have great physical and mental disabilities, with developmental skills usually stalled at the 6-month level. Major abnormalities include heart defects, a displaced liver, growth retardation, and oddly clenched fists. Milder signs include overlapping placement of fingers, a narrow



Figure 13.14 Trisomies 18 and 13. (a) An infant with trisomy 18 clenches the fists in an odd way, with fingers overlapping. (b) Very few babies with trisomy 13 are as healthy as Hazel. Most die in infancy.

and flat skull, abnormally shaped and low-set ears, a small mouth and face, unusual or absent fingerprints, short, large toes with fused second and third toes, and “rocker-bottom” feet. Most cases of trisomy 18 are traced to nondisjunction in meiosis II of the oocyte.

Trisomy 13—Tykesia’s Story

At 15 months of age, Tykesia is a “long-term survivor” of trisomy 13. About 92 percent of infants born with an extra chromosome 13 do not live to see their first birthdays.

Tykesia is small for her age, at the 5th percentile for weight, but she is happy, curious and playful. Her physical skills, however, lag. She can finally, with great effort, sit up, but cannot yet crawl. She has about 20 minor seizures a day, which look like jerks or startles, and has difficulty eating because of persistent acid reflux. She is also missing a rib. Early surgeries corrected a cleft lip and palate, removed an extra finger and toe, and corrected a hernia. Blood vessels leading from the heart to the lungs that did not close as they normally should before birth did so by the time Tykesia was 6 months old. She is mentally retarded, but her parents hope she will live long enough to attend preschool. Despite these challenges, Tykesia’s case is mild—she has her sight and hearing, unlike many others with trisomy 13.

Trisomy 13 has a different set of signs and symptoms than trisomy 18. Most striking is a fusion of the developing eyes, so that a fetus has one large eyelike structure in the center of the face. More common is a small or absent eye. Major abnormalities affect the heart, kidneys, brain, face, and limbs. The nose is often malformed, and cleft lip and/or palate is present in a small head. There may be extra fingers and toes. Ultrasound examination of an affected newborn often reveals more extensive anomalies than were visualized prenatally, such as an extra spleen, abnormal liver, rotated intestines, and an abnormal pancreas. A few individuals have survived until adulthood, but they do not progress developmentally beyond the 6-month level.

Sex Chromosome Aneuploids: Female

People with sex chromosome aneuploidy have extra or missing sex chromosomes. **Table 13.6** indicates how these aneuploids can arise. Note that some conditions can result from nondisjunction in meiosis in the male or female. These conditions are generally much less serious than autosomal aneuploids, which is why this book has no photos of affected individuals—they look like anyone else.

XO Syndrome—Miranda’s Story

Miranda was well into her teen years, but still looked about 12. It wasn’t just that she was short. Her breasts had never developed, and she still hadn’t gotten her first menstrual period. Her sister Charlotte, 2 years younger, actually looked older. When Miranda turned 16, her physician suggested that she have her chromosomes checked. While reassuring Miranda that delayed puberty could be treated, the doctor also explained what she was looking to rule out—absence of an X chromosome, called Turner or XO syndrome.

Table 13.6 How Nondisjunction Leads to Sex Chromosome Aneuploids

Situation	Oocyte	Sperm	Consequence
Normal	X	Y	46,XY normal male
	X	X	46,XX normal female
Female nondisjunction	XX	Y	47,XXY Klinefelter syndrome
	XX	X	47,XXX triplo-X
		Y	45,Y nonviable
		X	45,X Turner syndrome
Male nondisjunction (meiosis I)	X		45,X Turner syndrome
	X	XY	47,XXY Klinefelter syndrome
Male nondisjunction (meiosis II)	X	XX	47,XXX triplo-X
	X	YY	47,XYY Jacobs syndrome
	X		45,X Turner syndrome
Male and female nondisjunction	XX	YY	48,XXYY syndrome

Miranda indeed lacked a second X chromosome. The diagnosis explained other problems she’d had, some obvious, some rather subtle. She’d always had poor hearing and high blood pressure, and the syndrome also accounted for her low thyroid function and the “beauty marks” that dotted her skin. Life with a single X had also affected Miranda’s mind, although in mild ways—she had difficulty solving math problems that required envisioning objects in three-dimensional space, and had a poor memory. Miranda started taking estrogen and progesterone, which made her develop secondary sexual characteristics, and her doctor explained that one day she could use a donor egg to become pregnant. Had Miranda known about her unusual chromosomes before her teen years, she could have started taking growth hormones to maximize her height. But Miranda has no regrets—she’s smart and happy.

In 1938, at a medical conference, a U.S. endocrinologist named Henry Turner described seven young women, aged 15 to 23, who were sexually undeveloped, short, had folds of skin on the back of their necks, and had malformed elbows. (Eight years earlier, an English physician had described the syndrome in young girls, so it is called Ullrich syndrome in the U.K.) Alerted to what would become known as Turner syndrome in the United States, other physicians soon began identifying such patients. Physicians assumed that a hormonal insufficiency caused the symptoms. They were right, but there was more to the story—a chromosomal imbalance caused the hormone deficit.

In 1954, at a London hospital, a physician discovered that cells from Turner patients lacked a Barr body, the dark spot that indicates a second X chromosome. Might lack of a sex chromosome cause the symptoms, particularly failure to

mature sexually? By 1959, karyotyping confirmed the presence of only one X chromosome. Later, researchers learned that only 50 percent of affected individuals are XO. The rest are missing only part of an X chromosome or are mosaics, with only some cells missing an X.

Like the autosomal aneuploids, Turner syndrome, now called XO syndrome, is found more frequently among spontaneously aborted fetuses than among newborns—99 percent of XO fetuses are not born. The syndrome affects 1 in 2,500 female births. However, if amniocentesis or CVS was not done, a person with XO syndrome would likely not know she has a chromosome abnormality until she lags in sexual development. Two X chromosomes are necessary for normal sexual development in females.

At birth, a girl with XO syndrome looks normal, except for puffy hands and feet caused by impaired lymph flow. In childhood, signs of XO syndrome include wide-set nipples, soft nails that turn up at the tips, slight webbing at the back of the neck, short stature, coarse facial features, and a low hairline at the back of the head. About half of people with XO syndrome have impaired hearing and frequent ear infections due to a small defect in the shape of the coiled part of the inner ear. They cannot hear certain frequencies of sound.

At sexual maturity, sparse body hair develops, but the girls do not ovulate or menstruate, and their breasts do not develop. The uterus is very small, but the vagina and cervix are normal size. In the ovaries, oocytes speed through development, depleting the supply during infancy. Intelligence is normal. However, “Turner neurocognitive phenotype” may impair the ability to solve math problems that entail envisioning objects in three-dimensional space, and may cause memory deficits. Hormones (estrogen and progesterone) can be given to stimulate development of secondary sexual structures for individuals diagnosed before puberty, and prompt use of growth hormone can maximize height.

Individuals who are mosaics (only some cells lack the second X chromosome) may have children, but their offspring are at high risk of having abnormal numbers of chromosomes. XO syndrome is unrelated to the age of the mother. The effects of XO syndrome continue past the reproductive years. Life span is shortened slightly. Adults are more likely to develop certain disorders than the general population, including osteoporosis, types 1 and 2 diabetes, and colon cancer.

The many signs and symptoms of XO syndrome result from the loss of specific genes. For example, loss of a gonadal dysgenesis gene accounts for the ovarian failure, whereas absence of a transcription factor causes short stature. Deletion of another gene causes the hearing defect.

Triplo-X

About 1 in every 1,000 females has an extra X chromosome in each of her cells, a condition called triplo-X. The only symptoms are tall stature and menstrual irregularities. Although triplo-X females are rarely mentally retarded, they tend to be less intelligent than their siblings. The lack of symptoms reflects the protective effect of X inactivation—all but one of the X chromosomes is inactivated.

Sex Chromosome Aneuploids: Male

Any individual with a Y chromosome is a male.

XXY Syndrome—Stefan’s Story

Looking back, Stefan Schwarz’s only indication of XXY syndrome was small testes. When his extra X chromosome was detected when he was 25, suddenly his personality quirks made sense.

“I was very shy, reserved, and had trouble making friends. I would fly into rages for no apparent reason. My parents knew when I was very young that there was something about me that wasn’t right,” he recalls.

Many psychologists, psychiatrists, and therapists diagnosed “learning disabilities,” and one even told Stefan he “was stupid and lazy, and would never amount to anything.” But Stefan proved them wrong. He earned two bachelor’s degrees, then started a successful career as a software engineer. Today he heads a support group for men with XXY syndrome.

About 1 in 500 males has the extra X chromosome that causes XXY (Klinefelter) syndrome. Severely affected men are underdeveloped sexually, with rudimentary testes and prostate glands and sparse pubic and facial hair. They have very long arms and legs, large hands and feet, and may develop breast tissue. XXY syndrome is the most common genetic or chromosomal cause of male infertility.

Testosterone injections during adolescence can limit limb lengthening and stimulate development of secondary sexual characteristics. Boys and men with XXY syndrome may be slow to learn, but they are usually not mentally retarded unless they have more than two X chromosomes, which is rare.

Men with XXY syndrome have fathered children, with medical assistance. Doctors select sperm that contain only one sex chromosome and use the sperm to fertilize oocytes. However, sperm from men with XXY syndrome are more likely to have extra chromosomes—usually X or Y, but also autosomes—than sperm from men who do not have XXY syndrome.

XXYY Syndrome—Devon’s Story

Devon’s parents suspected early on that he was different. His problems were so common, however, that it was years before a chromosome check revealed an extra X and an extra Y.

As a toddler, Devon sat, crawled, and walked more slowly than normal. He was also late to talk, and when he did, his speech was not easily understood. In preschool, problems emerged. He’d have frequent outbursts that seemed to come out of nowhere, and would make inappropriate comments, such as telling a classmate her dress was ugly. He was tall and clumsy, and drooled and choked easily. Devon would run about flapping his arms, then hide under a chair. Severe ulcers formed on his legs. These problems made it difficult for him to make friends.

By the second grade, Devon’s difficulties alarmed his special education teacher, who suggested to Devon’s parents, Drucilla and Neil, that they have his chromosomes checked. They did. Since Drucilla and Neil’s chromosomes were normal, Devon must have been conceived from a very unusual oocyte.

meeting a very unusual sperm, both arising from nondisjunction. The extra sex chromosomes explained nearly all of the boy's problems, and even a few that hadn't been recognized, such as curved pinkies, flat feet, and scoliosis. He began receiving testosterone injections so that his teen years would be more normal than his difficult childhood had been.

A male with an extra X chromosome and an extra Y chromosome was until recently classified as having Klinefelter syndrome. Increased use of amniocentesis to check fetal chromosomes, however, has made it possible to distinguish XYY from XXY individuals. Even though they share many characteristics, those with the second Y have more severe behavioral problems and tend to develop foot and leg ulcers, resulting from poor venous circulation.

In XYY syndrome, childhood and adolescence often include attention deficit disorder, obsessive compulsive disorder, and learning disabilities. In the teen years, testosterone level is low, development of secondary sexual characteristics is delayed, and the testes are undescended. A man with XYY syndrome is infertile.

XYY Syndrome

In 1961, a tall, healthy man, known for his boisterous behavior, had a chromosome check after fathering a child with Down syndrome. The man had an extra Y chromosome. A few other cases were detected over the next several years.

In 1965, researcher Patricia Jacobs published results of a survey among 197 inmates at Carstairs, a high-security prison in Scotland. Of twelve men with unusual chromosomes, seven had an extra Y. Might their violent or aggressive behavior be linked to their extra Y chromosome? Jacobs's findings were repeated for mental institutions, and soon after, Newsweek magazine ran a cover story on "congenital criminals." Having an extra Y, known as Jacobs syndrome, became a legal defense for committing a violent crime.

In the early 1970s, newborn screens began in hospital nurseries in England, Canada, Denmark, and Boston. Social workers and psychologists visited XYY boys and offered "anticipatory guidance" to the parents on how to deal with their toddling future criminals. By 1974, geneticists and others halted the program, pointing out that singling out these boys on the basis of a few statistical studies was inviting self-fulfilling prophecy.

One male in 1,000 has an extra Y chromosome. Today, we know that 96 percent of XYY males are apparently normal. The only symptoms attributable to the extra chromosome may be great height, acne, and perhaps speech and reading problems. An explanation for the continued prevalence of XYY among mental-penal institution populations may be more psychological than biological. Large body size may lead teachers, employers, parents, and others to expect more of these people, and a few XYY individuals may deal with this stress aggressively.

Jacobs syndrome can arise from nondisjunction in the male, producing a sperm with two Y chromosomes that fertilizes an X-bearing oocyte. Geneticists have never observed a sex chromosome constitution of one Y and no X. Since the Y chromosome carries little genetic material, and the gene-packed

X chromosome would not be present, the absence of so many genes makes development beyond a few cell divisions in a YO embryo impossible.

Key Concepts

1. Polyploids have extra sets of chromosomes and do not survive for long.
2. Aneuploids have extra or missing chromosomes. Nondisjunction during meiosis causes aneuploidy.
3. Trisomies are less severe than monosomies, and sex chromosome aneuploidy is less severe than autosomal aneuploidy.
4. Mitotic nondisjunction produces chromosomal mosaics.
5. Down syndrome (trisomy 21) is the most common autosomal aneuploid, followed by trisomies 18 and 13.
6. Sex chromosome aneuploid conditions include XO, triplo-X, XXY, XYY, and XYY syndromes.

13.4 Abnormal Chromosome Structure

Structural chromosomal defects include missing, extra, or inverted genetic material within a chromosome or combined or exchanged parts of nonhomologs (translocations) (figure 13.15). Abnormal chromosomes are balanced if the normal amount of genetic material persists (inversions and balanced translocations) and unbalanced if excess or deficient DNA results (duplications and deletions, which may be caused by inversions or translocations).

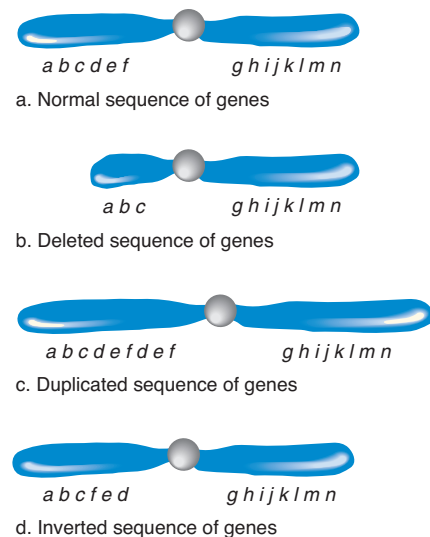


Figure 13.15 Chromosome abnormalities. If a hypothetical normal gene sequence appears as shown in (a), then (b) represents a deletion, (c) a duplication, and (d) an inversion.

Deletions and Duplications

Ashley's Story

Our daughter, Ashley Elizabeth Naylor (**figure 13.16**), was born August 12, 1988. The doctors suspected complications. Two weeks after her birth, chromosome analysis revealed *cri-du-chat* (cat cry) syndrome, also known as $5p^-$ syndrome because part of the short arm of one copy of chromosome 5 is missing. This is a rare disorder; we were told, and little could be offered to help our daughter. The doctors used the words “profoundly retarded.”

Ashley defied all the standard medical labels, as well as her doctors' expectations. Her spirit and determination enabled her to walk with the aid of a walker and express herself using sign language and a communication device. With early intervention and education, Ashley found the resources and additional encouragement she needed to succeed. In May of 1994, Ashley's small body could no longer support the spirit that inspired so many. She passed away after a long battle with pneumonia. Her physical presence is gone, but her message remains: hope.

Deletions and duplications are missing and extra DNA sequences, respectively. They are types of copy number variants (CNVs), introduced in chapter 12. Traditionally, only large deletions and duplications could be visualized in karyotypes and, in general, the more genes involved, the more severe the associated syndrome. Ashley Naylor was missing a sizable portion of her chromosome 5. **Figure 13.17** depicts a common duplication, of part of chromosome 15. Deletions and duplications



Figure 13.16 Ashley Naylor brought great joy to her family and community during her short life. She had $5p^-$ syndrome.

Courtesy of Kathy Naylor.

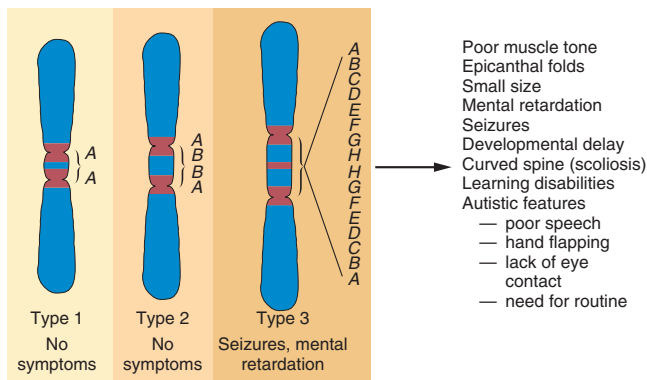


Figure 13.17 A duplication. A study of duplications of parts of chromosome 15 revealed that small duplications do not affect the phenotype, but larger ones may. The letters indicate specific DNA sequences, which serve as markers to compare chromosome regions. Note that the duplication is also inverted.

often arise “*de novo*,” which means that neither parent has the abnormality, and it is therefore new. In these cases, the symptoms can be attributed to the chromosomal abnormality.

A technique called comparative genomic hybridization is used to detect very small CNVs, which are also termed microdeletions and microduplications. The technique compares the abundance of copies of a particular CNV in the same amount of DNA from two sources—one with a medical condition, one healthy. Comparative genomic hybridization is being used increasingly to help narrow down diagnoses for children with autism, mental retardation, learning disabilities, or just strange behavior. For example, the technique showed that a young boy who had difficulty concentrating and sleeping and would often scream for no reason had a small duplication in chromosome 7. A young girl plagued with head-banging behavior, digestive difficulties, severe constipation, and great sensitivity to sound had a microdeletion in chromosome 16. Other microdeletions cause male infertility.

Deletions and duplications can arise from chromosome rearrangements. These include translocations, inversions, and ring chromosomes.

Translocation Down Syndrome

Rhiannon's Story

When Rhiannon P. was born, while her parents marveled at her beauty, the obstetrician was disturbed by her facial features: the broad, tilted eyes and sunken nose looked very slightly like the face of a child with Down syndrome. The doctor might not have noticed, except that the mother, Felicia, had had two spontaneous abortions. The family history suggested a chromosome problem. So when he examined the newborn, the doctor looked for the telltale single crease in the palms of people with Down syndrome, and found it. Gently, he told Felicia and her husband Matt that he'd like to do a chromosome check.

Two days later, the new parents learned that their daughter had an unusual form of Down syndrome that was inherited from

one of them, rather than the “extra chromosome” form of the condition that causes most cases. Since Matt’s mother and sister had also had several miscarriages, the suspected exchanged chromosomes likely came from his side. Karyotypes of Matt and Felicia confirmed this: Matt was a translocation carrier. One of his chromosome 14s had attached to one of his chromosome 21s, and distribution of the unusual chromosome in meiosis had led to various imbalances, depicted in **figure 13.18**.

Rhiannon had very mild Down syndrome. She did not have any of the physical problems associated with the condition, and she did well in school with the help of a special education teacher. Matt and Felicia chose to see the bright side—each conception would have a one in three chance of having balanced chromosomes. Some day they would give Rhiannon a brother or sister.

In a translocation, different (nonhomologous) chromosomes exchange or combine parts. Translocations can be inherited because they can be present in carriers, who have the normal amount of genetic material, but it is rearranged. A translocation can affect the phenotype if it breaks a gene

or leads to duplications or deletions in the chromosomes of offspring.

There are two major types of translocations, as well as rarer types. In a **Robertsonian translocation**, the short arms of two different acrocentric chromosomes break, leaving sticky ends on the two long arms that join, forming a single, large chromosome with two long arms (chromosome 14/21 in figure 13.18). The tiny short arms are lost, but their DNA sequences are repeated elsewhere in the genome, so the loss does not cause symptoms. The person with the large, translocated chromosome, called a **translocation carrier**, has 45 chromosomes, but may not have symptoms if no crucial genes have been deleted or damaged. Even so, he or she may produce unbalanced gametes—sperm or oocytes with too many or too few genes. This can lead to spontaneous abortion or birth defects.

In 1 in 20 cases of Down syndrome, a parent has a Robertsonian translocation between chromosome 21 and another, usually chromosome 14. That parent produces some gametes that lack either of the involved chromosomes and some gametes

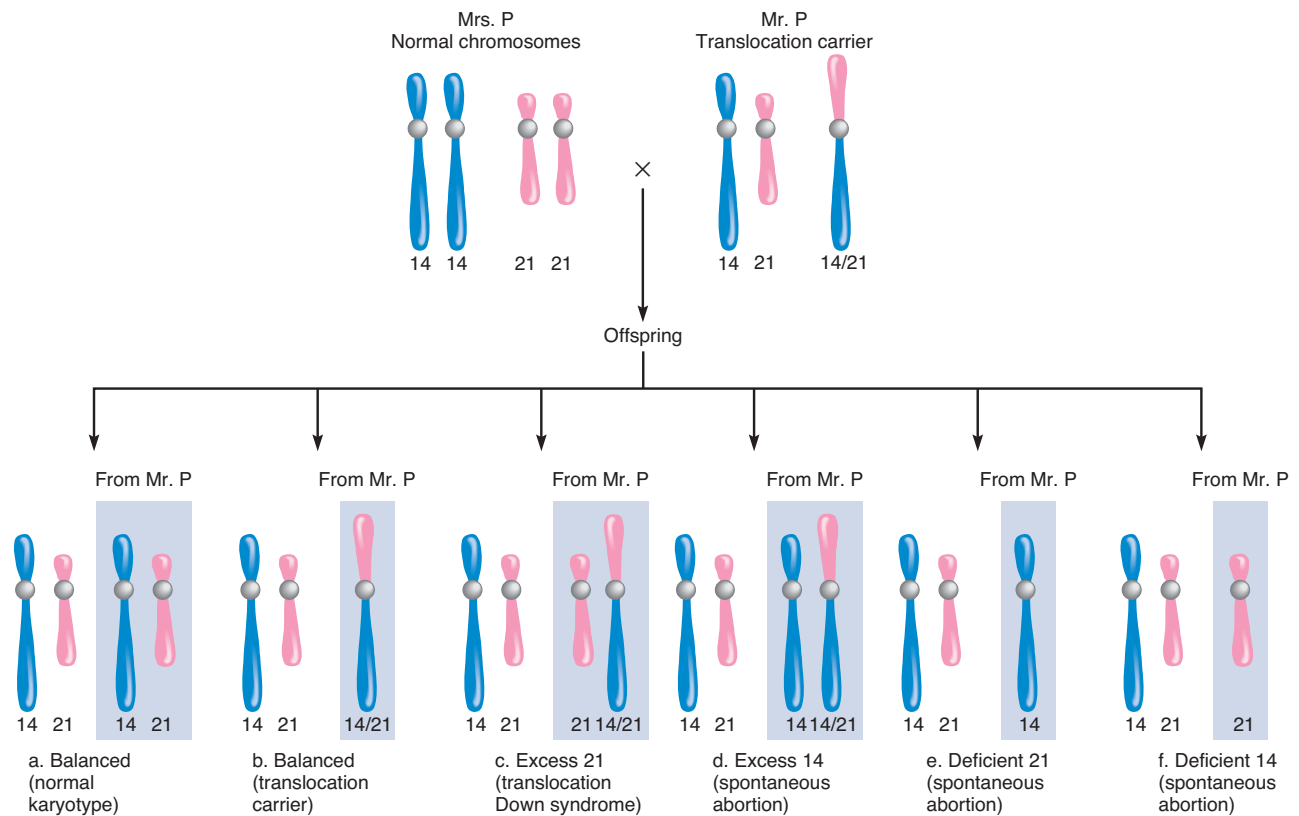


Figure 13.18 A Robertsonian translocation. Mr. P. has only 45 chromosomes because the long arm of one chromosome 14 has joined the long arm of one chromosome 21. He has no symptoms. Mr. P. makes six types of sperm cells, and they determine the fates of offspring. **(a)** A sperm with one normal chromosome 14 and one normal 21 yields a normal child. **(b)** A sperm carrying the translocated chromosome produces a child who is a translocation carrier, like Mr. P. **(c)** If a sperm contains Mr. P’s normal 21 and his translocated chromosome, the child receives too much chromosome 21 material and has Down syndrome. **(d)** A sperm containing the translocated chromosome and a normal 14 leads to excess chromosome 14 material, which is lethal in the embryo or fetus. If a sperm lacks either chromosome 21 **(e)** or 14 **(f)**, it leads to monosomies, which are lethal prenatally. (Chromosome arm lengths are not precisely accurate.)

that have extra material from one of the translocated chromosomes. In such a case, each fertilized ovum has a 1 in 2 chance of ending in spontaneous abortion, and a 1 in 6 chance of developing into an individual with Down syndrome. The risk of giving birth to a child with Down syndrome is theoretically 1 in 3, because the spontaneous abortions are not births. However, because some Down syndrome fetuses spontaneously abort, the actual risk of a couple in this situation having a child with Down syndrome is about 15 percent. The other two outcomes—a fetus with normal chromosomes or a translocation carrier like the parent—have normal phenotypes. Either a male or a female can be a translocation carrier, and the condition is not related to age.

Because Robertsonian translocations are among the more common chromosomal aberrations, an intriguing idea has arisen—they could one day lead to a human karyotype of 44 instead of 46 chromosomes, and perhaps even two types of people! Individuals who have one Robertsonian translocation have 45 chromosomes, and therefore may make gametes missing a chromosome, which impairs fertility. A person who has two different Robertsonian translocations would have 44 chromosomes, but the normal amount of genetic material. Two such people could have children together, the male producing sperm and the female producing oocytes with 22 chromosomes each. Robertsonian translocations affect one in 1,000 individuals. The chance of two people with different single translocations passing both to shared offspring is about 1 in 4 million—unlikely, yet possible.

In the second major type of translocation, a **reciprocal translocation**, two different chromosomes exchange parts (**figure 13.19**). About 1 in 500 people is a carrier for a reciprocal translocation. FISH can be used to highlight the involved chromosomes. If the chromosome exchange does not break any genes, then a person who has both translocated chromosomes is healthy and a translocation carrier. He or she has the normal amount of genetic material, but it is rearranged. A reciprocal translocation carrier can have symptoms if one of the two breakpoints lies in a gene, disrupting its function. Sometimes, a *de novo* translocation arises in a gamete that leads to a new individual with a disorder, as opposed to inheriting a translocated chromosome from a parent who is a carrier.

A rare type of translocation is an insertional translocation. In this situation, part of one chromosome inserts into a nonhomologous chromosome. Symptoms may result if a vital gene is disrupted or if genetic material is lost or present in excess. The chapter opener describes an insertional translocation in a family.

A carrier of any type of translocation can produce some unbalanced gametes—sperm or oocytes that have deletions or duplications of some of the genes in the translocated chromosomes. The resulting phenotype depends upon the particular genes that the chromosomal rearrangement disrupts and

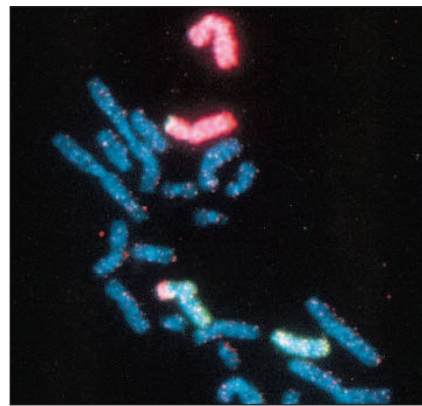
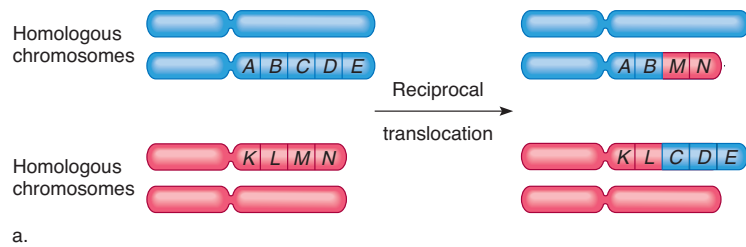


Figure 13.19 A reciprocal translocation. In a reciprocal translocation, two nonhomologous chromosomes exchange parts. In **(a)**, genes *C*, *D*, and *E* on the blue chromosome exchange positions with genes *M* and *N* on the red chromosome. Part **(b)** highlights a reciprocal translocation using FISH. The pink chromosome with the dab of blue, and the blue chromosome with a small section of pink, are the translocated chromosomes.

whether they are extra or missing. A translocation and a deletion can cause the same syndrome if they affect the same part of a chromosome.

A genetic counselor suspects a translocation when a family has a history of birth defects, pregnancy loss, and/or stillbirth. Prenatal testing may also reveal a translocation in a fetus, which can then be traced back to a parent who is a translocation carrier—or sometimes it is *de novo*.

Inversions

Madison's Story

Madison and Grant were excited about getting the results of the amniocentesis—Madison had never carried a pregnancy this far before. But they grew alarmed when the doctor's office called and asked them to come in to receive the results.

Prepared for the worst, the couple was surprised and confused to learn that the fetus had an inverted chromosome—some of the bands that normally appear on chromosome 11 were flipped around. What did it mean? Before the genetic counselor would give them information on genes that might be affected, she advised that the parents-to-be have their chromosomes checked. Although waiting another week for the results raised their anxiety level even higher, it was worth it, because Madison had the same inversion as the fetus! Because

Madison was healthy, the unusual chromosome would likely do their daughter no harm. When she was older, however, she might, like her mother, experience pregnancy loss.

An inverted sequence of chromosome bands is associated with health effects in only 5 to 10 percent of cases, in which the inversion disrupts important genes. If neither parent has the inversion, then it arose in a gamete. Effects may depend on which genes are involved. The human genome sequence can be consulted to identify genes that might be implicated in a particular inversion.

Like a translocation carrier, an adult heterozygous for an inversion can be healthy, but have reproductive problems. One woman had an inversion in the long arm of chromosome 15 and had two spontaneous abortions, two stillbirths, and two children with multiple problems who died within days of birth. She did eventually give birth to a healthy child. How did the inversion cause these problems?

Inversions with such devastating effects can be traced to meiosis, when a crossover occurs between the inverted chromosome segment and the noninverted homolog. To allow the genes to align, the inverted chromosome forms a loop. When crossovers occur within the loop, some areas are duplicated and some deleted in the resulting recombinant chromosomes. In inversions, the abnormal chromosomes result from the chromatids that crossed over.

Two types of inversions are distinguished by the position of the centromere relative to the inverted section. A **paracentric inversion** does not include the centromere (figure 13.20). A single crossover within the inverted segment gives rise to two normal and two very abnormal chromosomes. The other two chromosomes are normal. One abnormal chromosome retains both centromeres and is termed dicentric. When the cell divides, the two centromeres are pulled to opposite sides of the cell, and the chromosome breaks, leaving pieces with extra or missing segments. The second type of abnormal chromosome resulting from a crossover within an inversion loop is a small piece that lacks a centromere, called an acentric fragment. When the cell divides, the fragment is lost because a centromere is required for cell division.

A **pericentric inversion** includes the centromere within the loop. A crossover in it produces two chromosomes that have duplications and deletions, but one centromere each (figure 13.21).

Isochromosomes and Ring Chromosomes

Another meiotic error that leads to unbalanced genetic material is the formation of an isochromosome, which is a chromosome that has identical arms. This occurs when, during division, the centromeres part in the wrong plane (figure 13.22). Isochromosomes are known for chromosomes 12 and 21 and for the long arms of the X and the Y. Some women with Turner syndrome are not the more common XO, but have an isochromosome with the long arm of the X chromosome duplicated but the short arm absent.

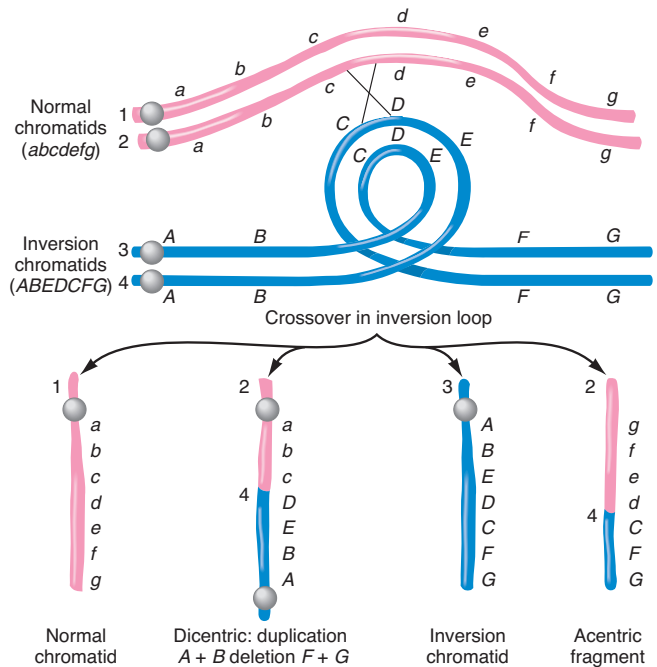


Figure 13.20 Paracentric inversion. A paracentric inversion in one chromosome leads to one normal chromatid, one inverted chromatid, one with two centromeres (dicentric), and one with no centromere (an acentric fragment) if a crossover occurs with the normal homolog. The letters *a* through *g* denote genes.

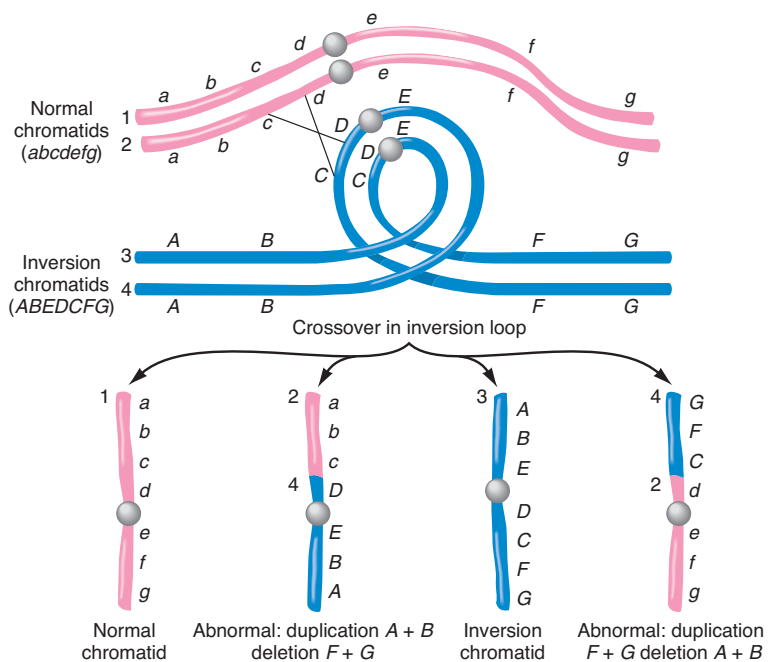


Figure 13.21 Pericentric inversion. A pericentric inversion in one chromosome leads to two chromatids with duplications and deletions, one normal chromatid, and one inverted chromatid if a crossover occurs with the normal homolog.

Chromosomes shaped like rings form in 1 out of 25,000 conceptions. Ring chromosomes may arise when telomeres are lost, leaving sticky ends that adhere. Exposure to radiation can also form rings. They can form from any chromosome, and may be present as one of the 46 chromosomes or in addition to them.

Most ring chromosomes consist of DNA repeats and do not affect health. Some do, however. This is the case for ring chromosome 20. When 6-year-old Cara Ford lost the ability to

walk, talk, or eat and developed seizures, no physician in her native United Kingdom could offer a diagnosis. Physicians at an epilepsy center in New York City detected the ring chromosome, and treated the seizures (**figure 13.23**). Cara's father, Stewart Ford, started the Ring Chromosome 20 Foundation to alert physicians to test patients with seizures for abnormal chromosomes.

Table 13.7 summarizes causes of different types of chromosomal aberrations.

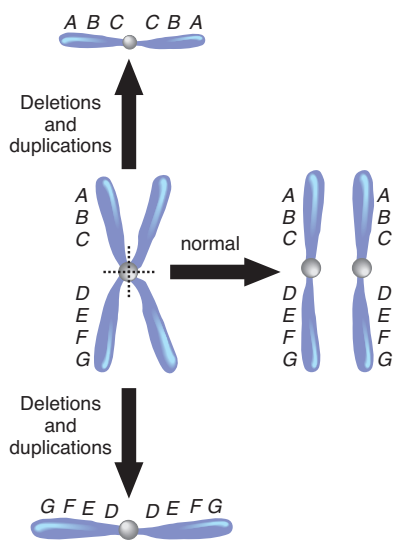


Figure 13.22 Isochromosomes have identical arms. They form when chromatids divide along the wrong plane (in this depiction, horizontally rather than vertically).

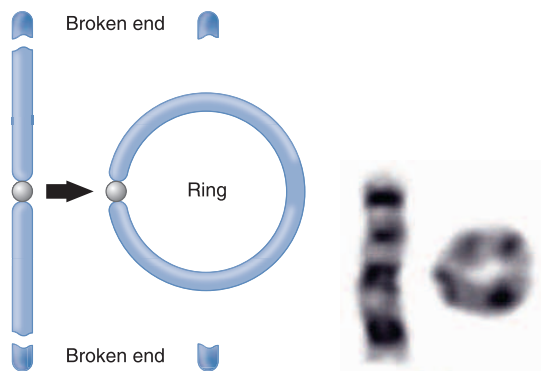


Figure 13.23 A ring chromosome. A ring chromosome may form if the chromosome's tips (telomeres) break, forming sticky ends. Genes can be lost or disrupted, possibly causing symptoms. Ring chromosome 20, for example, causes seizures.

Table 13.7 Causes of Chromosomal Aberrations

Abnormalities	Causes
Numerical Abnormalities	
Polyploidy	Error in cell division (meiosis or mitosis) in which not all chromatid pairs separate in anaphase
	Multiple fertilization
Aneuploidy	Nondisjunction (in meiosis or mitosis) leading to lost or extra chromosomes
Structural Abnormalities	
Deletions and duplications	Translocation
	Crossover between a chromosome that has a pericentric inversion and its noninverted homolog
Translocation	Exchange between nonhomologous chromosomes
Inversion	Breakage and reunion of fragment in same chromosome, but with wrong orientation
Dicentric and acentric	Crossover between a chromosome with a paracentric inversion and its noninverted homolog
Ring chromosome	A chromosome loses telomeres and the ends fuse, forming a circle

Key Concepts

1. Chromosome rearrangements can cause deletions and duplications.
2. In a Robertsonian translocation, the long arms of two different acrocentric chromosomes join.
3. In a reciprocal translocation, chromosomes exchange parts.
4. If a translocation leads to a deletion or duplication, or disrupts a gene, symptoms may result.
5. Gene duplications and deletions can occur in isochromosomes and ring chromosomes, and when crossovers involve inversions.
6. An isochromosome has two identical arms, introducing duplications and deletions.
7. Ring chromosomes form when telomeres are missing.

13.5 Uniparental Disomy—A Double Dose from One Parent

If nondisjunction occurs in sperm and oocyte, a pair of chromosomes (or their parts) can come solely from one parent, rather than one from each parent, as Mendel's law of segregation predicts. For example, if a sperm lacking a chromosome 14 fertilizes an ovum with two copies of that chromosome, an individual with the normal 46 chromosomes results, but the two chromosome 14s come only from the female.

Inheriting two chromosomes or chromosome segments from one parent is called **uniparental disomy** (UPD) ("two bodies from one parent"). UPD can also arise from a trisomic embryo in which some cells lose the extra chromosome, leaving two homologs from one parent. For example, an embryo may have trisomy 21, with the extra chromosome 21 coming from the father. If in some cells the chromosome 21 from the mother is lost, then both remaining copies of the chromosome are from the father.

Because UPD requires the simultaneous occurrence of two very rare events—either nondisjunction of the same chromosome in sperm and oocyte, or trisomy followed by chromosome loss—it is very rare. In addition, many cases are probably never seen, because bringing together identical homologs inherited from one parent could give the fertilized ovum a homozygous set of lethal alleles. Development would halt. Other cases of UPD may go undetected if they cause known recessive conditions and both parents are assumed to be carriers, when actually only one parent contributed to the offspring's illness. This was how UPD was discovered.

In 1988, Arthur Beaudet of the Baylor College of Medicine saw a very unusual patient with cystic fibrosis. Beaudet was comparing CF alleles of the patient to those of her parents, and he found that only the mother was a carrier—the father had two normal alleles. Beaudet constructed haplotypes for each parent's chromosome 7, which includes the CF gene, and he found that the daughter had two copies from her mother, and none from her father (**figure 13.24**). How did this happen?

Apparently, in the patient's mother, nondisjunction of chromosome 7 in meiosis II led to formation of an oocyte bearing two identical copies of the chromosome, instead of the usual one. A sperm that had also undergone nondisjunction and lacked a chromosome 7 then fertilized the abnormal oocyte. The mother's extra genetic material compensated for the father's deficit, but unfortunately, the child inherited a double dose of the mother's chromosome that carried the mutant CF allele. In effect, inheriting two of the same chromosome from one parent shatters the protection that combining genetic material from two individuals offers, a protection that is the defining characteristic of sexual reproduction.

UPD may also cause disease if it removes the contribution of the important parent for an imprinted gene. Recall from chapter 6 that an imprinted gene is expressed if it comes from one parent, but silenced if it comes from the other (see figure 6.15). If UPD removes the parental genetic material

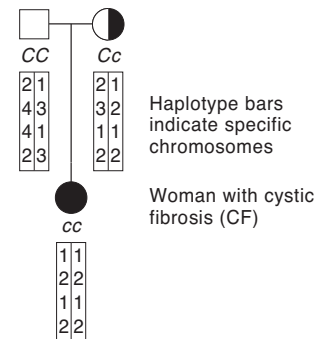


Figure 13.24 Uniparental disomy. Uniparental disomy doubles part of one parent's genetic contribution. In this family, the woman with CF inherited two copies of her mother's chromosome 7, and neither of her father's. Unfortunately, it was the chromosome with the disease-causing allele that she inherited in a double dose.

that must be present for a critical gene to be expressed, a mutant phenotype results. The classic example is the 20 to 30 percent of Prader-Willi syndrome and Angelman syndrome cases caused by UPD (see figure 6.16). These disorders arise from mutations in different genes that are closely linked in a region of the long arm of chromosome 15, where imprinting occurs. They both cause mental retardation and a variety of other symptoms, but are quite distinct.

In 1989, researchers found that some children with Prader-Willi syndrome have two parts of the long arm of chromosome 15 from their mothers. The disease results because the father's Prader-Willi gene must be expressed for the child to avoid the associated illness. For Angelman syndrome, the situation is reversed. Children have a double dose of their father's DNA in the same chromosomal region implicated in Prader-Willi syndrome, with no maternal contribution. The mother's gene must be present for health.

People usually learn their chromosomal makeup only when something goes wrong—when they have a family history of reproductive problems, exposure to a toxin, cancer, or symptoms of a known chromosomal disorder. While researchers analyze the human genome sequence, chromosome studies will continue to be part of medical care—beginning before birth.

Key Concepts

1. Uniparental disomy (UPD) results when two chromosomes or chromosome parts are inherited from the same parent.
2. It can arise from two nondisjunction events or a trisomy and subsequent chromosome loss.
3. UPD can cause disease if it creates a homozygous recessive condition, or if it disrupts imprinting.

Summary

13.1 Portrait of a Chromosome

1. Mutation can occur at the chromosomal level. **Cytogenetics** is the study of chromosome aberrations and their effects on phenotypes.
2. **Heterochromatin** stains darkly and harbors many DNA repeats. **Euchromatin** is light staining and contains many protein-encoding genes.
3. A chromosome consists of DNA and proteins. Essential parts are the **telomeres**, **centromeres**, and origin of replication sites.
4. Centromeres include DNA repeats and proteins that enable the cell to divide.
5. Subtelomeres have telomere-like repeats that gradually change inward toward the centromere, as protein-encoding genes predominate.
6. Chromosomes are distinguishable by size, centromere position, satellites, and staining patterns. They are displayed in **karyotypes**.
7. A **metacentric** chromosome has two fairly equal arms. A **submetacentric** chromosome has a large arm and a short arm. An **acrocentric** chromosome's centromere is near a tip, so that it has one long arm and one very short arm.

13.2 Visualizing Chromosomes

8. Chromosomes can be obtained from any cell that has a nucleus. Prenatal diagnostic techniques include **amniocentesis**, **chorionic villus sampling**, **chromosome microarray analysis**, and fetal cell and nucleic acid detection. Serum markers screen fetuses at increased risk for chromosomal abnormalities.
9. Fluorescence *in situ* hybridization provides more specific chromosome bands than dyes. Ideograms display chromosome bands.
10. Chromosomal shorthand indicates chromosome number, sex chromosome constitution, and type of abnormality.

13.3 Abnormal Chromosome Number

11. A **euploid** somatic human cell has 22 pairs of autosomes and one pair of sex chromosomes.
12. **Polyloid** cells have extra chromosome sets.
13. **Aneuploids** have extra or missing chromosomes. **Trisomies** (an extra chromosome) are less harmful than **monosomies**

(lack of a chromosome), and sex chromosome aneuploidy is less severe than autosomal aneuploidy. **Nondisjunction** is uneven distribution of chromosomes in meiosis. It causes aneuploidy. Most autosomal aneuploids cease developing as embryos.

13.4 Abnormal Chromosome Structure

14. **Deletions** and/or **duplications** result from crossing over after pairing errors in synapsis. Crossing over in an inversion heterozygote can also generate deletions and duplications. Microdeletions and microduplications explain many disorders.
15. In a **Robertsonian translocation**, the short arms of two acrocentric chromosomes break, leaving sticky ends on the long arms that join to form an unusual, large chromosome.
16. In a **reciprocal translocation**, two nonhomologous chromosomes exchange parts.
17. An insertional translocation places a DNA sequence from one chromosome into a nonhomologous chromosome.
18. A **translocation carrier** may have an associated phenotype and produces some unbalanced gametes.
19. A heterozygote for an inversion may have reproductive problems if a crossover occurs between the inverted region and the noninverted homolog, generating deletions and duplications. A **paracentric inversion** does not include the centromere; a **pericentric inversion** does.
20. Isochromosomes repeat one chromosome arm but delete the other. They form when the centromere divides in the wrong plane during meiosis. Ring chromosomes form when telomeres are removed, leaving sticky ends that adhere.

13.5 Uniparental Disomy—A Double Dose from One Parent

21. In **uniparental disomy**, a chromosome, or a part of one, doubly represents one parent. It can result from nondisjunction in both gametes, or from a trisomic cell that loses a chromosome, leaving two from the same parent.
22. Uniparental disomy causes symptoms if it creates a homozygous recessive state associated with an illness, or if it affects an imprinted gene.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

1. What are the essential components of a chromosome? Of a centromere?
2. Which parts of chromosome structure are probably important in an evolutionary sense? What is the evidence for this?
3. How does the DNA sequence change with distance from the telomere?
4. How are centromeres and telomeres alike?

5. Distinguish among a euploid, aneuploid, and polyploid.
6. What happens during meiosis to produce
 - a. an aneuploid?
 - b. a polyploid?
 - c. the increased risk of trisomy 21 Down syndrome in the offspring of a woman over age 40 at the time of conception?
 - d. recurrent spontaneous abortions to a couple in which the man has a pericentric inversion?
 - e. several children with Down syndrome in a family where one parent is a translocation carrier?
7. A human liver has patches of cells that are octaploid—that is, they have eight sets of chromosomes. Explain how this might arise.
8. Describe an individual with each of the following chromosome constitutions. Mention the person's sex and possible phenotype.
 - a. 47,XXX
 - b. 45,X
 - c. 47,XX, trisomy 21
9. Which chromosomal anomaly might you expect to find more frequently among the members of the National Basketball Association than in the general population? Cite a reason for your answer.
10. About 80 percent of cases of Edward syndrome are caused by trisomy 18; 10 percent are caused by mosaic trisomy 18, and 10 percent are attributed to translocation. Distinguish among these three chromosome aberrations.
11. List three examples illustrating the idea that the amount of genetic material involved in a chromosomal aberration affects the severity of the associated phenotype.
12. List three types of chromosomal aberrations that can cause duplications and/or deletions, and explain how they do so.
13. Distinguish among three types of translocations.
14. Why would having the same inversion on both members of a homologous chromosome pair *not* lead to unbalanced gametes, as having the inversion on only one chromosome would?
15. Define or describe the following technologies:
 - a. FISH
 - b. amniocentesis
 - c. chorionic villus sampling
 - d. fetal cell sorting
16. Why are trisomies 13 and 18 more common at birth than trisomies 5 or 16?
17. How many chromosomes would a person have who has Klinefelter syndrome and also trisomy 21?
18. Explain why a female cannot have XXY syndrome and a male cannot have XO syndrome.
19. List three causes of Turner syndrome.

Applied Questions

1. Identify the structures and/or DNA sequences that must be present for a chromosome to carry information and withstand the forces of cell division.
2. Amniocentesis indicates that a fetus has the chromosomal constitution 46, XX, del(5)(p15). What does this mean? What might the child's phenotype be?
3. What type of test could determine whether a triploid infant resulted from a diploid oocyte fertilized by a haploid sperm, or from two sperm fertilizing one oocyte?
4. For an exercise in a college genetics laboratory course, a healthy student constructs a karyotype from a cell from the inside of her cheek. She finds only one chromosome 3 and one chromosome 21, plus two unusual chromosomes that do not seem to have matching partners.
 - a. What type of chromosomal abnormality does she have?
 - b. Why doesn't she have any symptoms?
 - c. Would you expect any of her relatives to have any particular medical problems? If so, which medical conditions?
5. A fetus ceases developing in the uterus. Several of its cells are karyotyped. Approximately 75 percent of the cells are diploid, and 25 percent are tetraploid (four copies of each chromosome). What do you think happened? When in development did it probably occur?
6. Distinguish among Down syndrome caused by aneuploidy, mosaicism, and translocation.
7. A couple has a son diagnosed with XXY syndrome. Explain how the son's chromosome constitution could have arisen from either parent.
8. DiGeorge syndrome (MIM 188400) causes abnormal parathyroid glands, heart defect, and an underdeveloped thymus gland. About 85 percent of patients have a microdeletion of part of chromosome 22. A girl, her mother, and a maternal aunt have very mild DiGeorge syndrome. They all have a reciprocal translocation of chromosomes 22 and 2.
 - a. How can a microdeletion and a translocation cause the same symptoms?
 - b. Why were the people with the translocation less severely affected than the people with the microdeletion?
 - c. What other problems might arise in the family with the translocation?
9. From 2 to 6 percent of people with autism have an extra chromosome that consists of two long arms of chromosome 15. The unusual chromosome includes two copies of the chromosome 15 centromere. Two normal copies of the chromosome are also present. What type of chromosome abnormality in a gamete can lead to this karyotype, which is called isodicentric 15?

Web Activities

10. Go to one of the websites listed in table 13.3, or find a similar disease organization, and learn about daily life with a particular chromosomal abnormality. Identify a challenge or problem common to several chromosomal syndromes, and describe how families cope with the problem.
11. Go to the website for the Baylor College of Medicine Medical Genetics Laboratories (<http://www.bcm.edu/cma/index.htm>). Select "Reference Table," and then select "Abnormalities Detected."
 - a. Select an abnormality.
 - b. Describe the mutation at the chromosomal level.
 - c. Click on the MIM # on the left, go to MIM, and use the information to describe the disorder.
12. Go to the website for the Genetic Science Learning Center at the Eccles Institute of Human Genetics at the University of Utah. Follow the instructions to create a karyotype.
13. Visit the website for the Human Genome Landmarks poster. Select a chromosome, and use Mendelian Inheritance in Man (MIM) to describe four traits or disorders associated with it. Or, consult the website for the Human Chromosome Launchpad for information on four genes carried on a specific chromosome.

Case Studies and Research Results

14. An ultrasound of a pregnant woman detects a fetus and a similarly sized and shaped structure that has disorganized remnants of facial features at one end. Amniocentesis on both structures reveals that the fetus is 46,XX, but cells of the other structure are 47,XX, trisomy 2. No cases of trisomy 2 infants have ever been reported. However, individuals who are mosaics for trisomy 2 have a collection of defects, including a rotated and underdeveloped small intestine, a small head, a hole in the diaphragm, and seizures.
 - a. How do the chromosomes of cells from the fetus and the other structure differ?
 - b. What is the process that occurred during meiosis to yield the bizarre structure?
 - c. Use this information to explain why children with a complete extra chromosome 2 are not seen, even though people with an extra chromosome 21 can live many years.
 - d. List two factors that determine the type and severity of abnormalities in an individual who is an aneuploid mosaic.
15. Two sets of parents who have children with Down syndrome meet at a clinic. The Phelps know that their son has trisomy 21. The Watkins have two affected children, and Mrs. Watkins has had two spontaneous abortions. Why should the Watkins be more concerned about future reproductive problems than the Phelps? How are the offspring of the two families different, even though they have the same symptoms?
16. The genomes of four of 291 people with mental retardation have a microdeletion in chromosome 17q21.3. The children have large noses, delayed speech, and mild mental retardation. Each had a parent with an inversion in the same part of chromosome 17.
 - a. Which arm of chromosome 17 is implicated in this syndrome?
 - b. How can an inversion in a parent's chromosome cause a deletion in a child's chromosome?
 - c. What other type of chromosome abnormality might occur in these children's siblings?
17. A 38-year-old woman, Dasheen, has amniocentesis. She learns that the fetus she is carrying has an inversion in chromosome 9 and a duplication in chromosome 18. She and her husband Franco have their chromosomes tested, and they learn that she has the duplication and Franco has the inversion. Both of the parents are healthy. Should they be concerned about the health of the fetus? Cite a reason for your answer.
18. Return to the chapter opener.
 - a. What type of translocation do Esteban and Maribella have?
 - b. Why doesn't their father have symptoms resulting from his abnormal chromosomes?
 - c. What are two possible chromosome configurations for Marcos?

PART 4 Population Genetics



A forensic scientist consults a DNA profile. The black bars represent short tandem repeats that form patterns used to exclude suspects in a crime.

CHAPTER

14

Constant Allele Frequencies

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- 14.1 Population Genetics Underlies Evolution**
- 14.2 Constant Allele Frequencies**
 - Hardy-Weinberg Equilibrium
 - Solving a Problem: The Hardy-Weinberg Equation
- 14.3 Applying Hardy-Weinberg Equilibrium**
- 14.4 DNA Profiling Uses Hardy-Weinberg Assumptions**
 - DNA Profiling Began with Forensics
 - Using Population Statistics to Interpret DNA Profiles
 - Using DNA Profiling to Identify Victims
 - Genetic Privacy

The Innocence Project Uses DNA Testing to Overturn Convictions

Josiah Sutton had served 4 1/2 years of a 25-year sentence for rape when he was exonerated, thanks to the Innocence Project. The nonprofit legal clinic and public policy organization created in 1992 has so far used DNA retesting to free more than 240 wrongfully convicted prisoners, most of whom were “poor, forgotten, and have used up all legal avenues for relief,” according to the website (www.innocenceproject.com). Sutton became a suspect after a woman in Houston identified him and a friend on the street 5 days after she had been raped, threatened with a gun, and left in a field. The two young men supplied saliva and blood samples, from which DNA profiles were done and compared to DNA profiles from semen found in the victim and in her car. At the trial, a crime lab employee testified that the probability that Sutton’s DNA matched that of the evidence by chance was 1 in 694,000—which led to his conviction. Jurors ignored the fact that Sutton’s physical description did not match the victim’s description of her assailant.

The DNA evidence came from more than one individual, yielded different results when the testing was repeated, and most importantly, looked at only seven of the parts of the genome that are typically compared in a DNA profile, or fingerprint. Doing the test correctly revised the statistics dramatically: Sutton’s pattern was shared not with 1 in 694,000 black men, as had originally been claimed, but with 1 in 16.

While in jail, Sutton read about DNA profiling and requested independent testing, but was refused. Then he got lucky. Journalists investigating the Houston crime laboratory became interested in his case, contacted a noted criminologist, and the Innocence Project became involved. Retesting the DNA evidence set him free. DNA profiling is a direct application of population genetics.

14.1 Population Genetics Underlies Evolution

Genetic change on an individual level takes the form of mutations and polymorphisms, the topics of chapters 12 and 13. Genetic change occurs in populations in the form of allele (gene variant) frequencies, which provide a different type of information than that yielded from DNA sequences. DNA profiling is a practical application of the study of genes in populations. Changing allele frequencies also underlie evolution, which is the basis of all biology.

A **population** is any group of members of the same species in a given geographical area who are potentially capable of mating and producing fertile offspring (**figure 14.1**). Examples of human populations are the students in a class, a stadium full of people, and the residents of a community, state, or nation. **Population genetics** is a branch of genetics that considers all the alleles in a population, which constitute the **gene pool**. The “pool” in gene pool refers to a collection of gametes, and an offspring represents two gametes from the pool. Alleles can move between populations when individuals migrate and mate. This movement, termed gene flow, underlies evolution, which is explored in the next two chapters.

It is at the population level that genetics goes beyond science, embracing information from history, anthropology, human behavior, and sociology. Population genetics enables us to trace our beginnings as well as understand our diversity today, and even predict the future.

Thinking about genes at the population level begins by considering frequencies—that is, how often a particular gene variant occurs in a particular population. Such frequencies can be calculated for alleles, genotypes, or phenotypes. For example, an allele frequency for the cystic fibrosis (CF) gene might be the number of $\Delta F508$ alleles among the residents of



Figure 14.1 A population is a group of organisms of the same species living in the same place. Populations of sexually reproducing organisms include many genetic variants. This genetic diversity gives the group a flexibility that enhances species survival. To us, these hippos look alike, but they can undoubtedly recognize phenotypic differences in each other.

San Francisco. $\Delta F508$ is the most common allele that, when homozygous, causes the disorder. The allele frequency derives from the two $\Delta F508$ alleles in each person with CF, plus those carried in heterozygotes, as a proportion of all alleles for that gene in the gene pool of San Francisco. The genotype frequencies are the proportions of heterozygotes and the two types of homozygotes in the population. Finally, a phenotypic frequency is simply the percentage of people in the population who have CF (or who do not). With multiple alleles for a single gene, the situation becomes more complex because there are many more phenotypes and genotypes to consider.

Phenotypic frequencies are determined empirically—that is, by observing how common a condition or trait is in a population. These figures have value in genetic counseling in estimating the risk that a particular inherited disorder will occur in an individual when there is no family history of the illness. **Table 14.1** shows disease incidence for phenylketonuria (PKU), an inborn error of metabolism that causes mental retardation unless the person follows a special, low-protein diet from birth. Note how the frequency differs in different populations.

On a broader level, shifting allele frequencies in populations reflect small steps of genetic change, called **microevolution**. These small, step-by-step changes underlie evolution. Genotype frequencies can change when any of the following conditions are met:

1. Individuals of one genotype are more likely to produce offspring with each other than with those of other genotypes (*nonrandom mating*).
2. Individuals *migrate* between populations.
3. Reproductively isolated small groups form within or separate from a larger population (*genetic drift*).
4. *Mutation* introduces new alleles into a population.
5. People with a particular genotype are more likely to produce viable, fertile offspring under a specific environmental condition than individuals with other genotypes (*natural selection*).

In today’s world, all of these conditions, except mutation, are quite common. Therefore, genetic equilibrium—when allele frequencies are *not* changing—is rare. Put another way,

Table 14.1 Frequency of PKU in Various Populations

Population	Frequency of PKU
Chinese	1/16,000
Irish, Scottish, Yemenite Jews	1/5,000
Japanese	1/119,000
Swedes	1/30,000
Turks	1/2,600
United States Caucasians	1/10,000

given our tendency to pick our own partners and move about, microevolution is not only possible, but also nearly unavoidable. (Chapter 15 considers these factors in depth.)

When enough microevolutionary changes accumulate to keep two fertile organisms of opposite sex in a population from successfully producing fertile offspring together, **macroevolution**, or the formation of a new species, has occurred. Before we consider the pervasive genetic evidence for evolution, this chapter discusses the interesting, but unusual, situation in which certain allele frequencies stay constant, a condition called **Hardy-Weinberg equilibrium**.

Key Concepts

1. Population genetics is the study of allele frequencies in groups of organisms of the same species in the same geographic area.
2. The genes in a population comprise its gene pool.
3. Microevolution reflects changes in allele frequencies in populations. It is not occurring if allele frequencies stay constant over generations (Hardy-Weinberg equilibrium).
4. Five factors can change genotype frequencies: nonrandom mating, migration, genetic drift, mutation, and natural selection.

14.2 Constant Allele Frequencies

Population genetics looks at phenotypes and genotypes among large numbers of individuals. Allele frequencies reveal the underlying rules. Tracking allele frequencies from one generation to the next can reveal evolution in action—or, if allele frequencies don't change, the state of Hardy-Weinberg equilibrium.

Hardy-Weinberg Equilibrium

In 1908, a Cambridge University mathematician named Godfrey Harold Hardy (1877–1947) and Wilhelm Weinberg (1862–1937), a German physician interested in genetics, independently used algebra to explain how allele frequencies can be used to predict phenotypic and genotypic frequencies in populations of diploid, sexually reproducing organisms.

Hardy unintentionally cofounded the field of population genetics with a simple letter published in the journal *Science*—he did not consider his idea to be worthy of the more prestigious British journal *Nature*. The letter began with a curious mix of modesty and condescension:

I am reluctant to intrude in a discussion concerning matters of which I have no expert knowledge, and I should have expected the very simple point which I wish to make to have been familiar to biologists.

Hardy continued to explain how mathematically inept biologists had incorrectly deduced from Mendel's work that

dominant traits would increase in populations while recessive traits would become rarer. At first glance, this seems logical. However, it is untrue for several reasons. Recessive alleles are introduced into a population by mutation or migration; maintained in heterozygotes; and become more common when they confer a reproductive advantage, thanks to natural selection.

Hardy and Weinberg disproved the assumption that dominant traits increase while recessive traits decrease using the language of algebra. The expression of population genetics in algebraic terms begins with the simple equation

$$p + q = 1.0$$

where p represents all dominant alleles for a gene, and q represents all recessive alleles. The expression " $p + q = 1.0$ " simply means that all the dominant alleles and all the recessive alleles comprise all the alleles for that gene in a population.

Next, Hardy and Weinberg described the possible genotypes for a gene with two alleles using the binomial expansion

$$p^2 + 2pq + q^2 = 1.0$$

In this equation, p^2 represents the percentage of homozygous dominant individuals, q^2 represents the percentage of homozygous recessive individuals, and $2pq$ represents the percentage of heterozygotes (**figure 14.2**). The letter p designates the frequency of a dominant allele, and q is the frequency of a recessive allele. **Figure 14.3** shows how the binomial expansion is derived from allele frequencies. Note that the derivation is conceptually the same as tracing alleles in a monohybrid cross.

p	+	q	=	1
All dominant alleles		All recessive alleles		Total number of alleles
p^2	+	$2pq$	+	q^2
Homozygous dominant		Heterozygous		Homozygous recessive
			=	1
				Total number of alleles for the gene in the population

Figure 14.2 The Hardy-Weinberg equation in English.

		Sperm	
		$p=A$	$q=a$
Oocytes	$p=A$	$p \times p = AA$	$p \times q = Aa$
	$q=a$	$p \times q = Aa$	$q \times q = aa$

Figure 14.3 Source of the Hardy-Weinberg equation. A variation on a Punnett square reveals how random mating in a population in which gene A has two alleles— A and a —generates genotypes aa , AA , and Aa , in the relationship $p^2 + 2pq + q^2$.

The binomial expansion used to describe genes in populations became known as the Hardy-Weinberg equation. It can reveal the changes in allele frequency that underlie evolution. If the proportion of genotypes remains the same from generation to generation, as the equation indicates, then that gene is not evolving (changing). This situation, Hardy-Weinberg equilibrium, is an idealized state. It is possible only if the population is large, if its members mate at random, and if no migration, genetic drift, mutation, or natural selection takes place.

Logic must be used with mathematical expressions to understand when Hardy-Weinberg equilibrium is occurring. It is rare for protein-encoding genes that affect the phenotype, because an organism's appearance and health affect its ability to reproduce. That is, genes that affect the phenotype are subject to natural selection—harmful allelic combinations are weeded out of the population. However, Hardy-Weinberg equilibrium is seen in DNA repeats and other sequences that do not affect the phenotype, and therefore are not subject to natural selection.

⊕ Solving a Problem

The Hardy-Weinberg Equation

We can follow the frequency of two alleles of a particular gene from one generation to the next to understand Hardy-Weinberg equilibrium. Mendel's laws underlie such population-based calculations.

Consider an autosomal recessive trait: a middle finger shorter than the second and fourth fingers. If we know the frequencies of the dominant and recessive alleles, then we can calculate the frequencies of the genotypes and phenotypes and trace the trait through the next generation. The dominant allele D confers normal-length fingers; the recessive allele d confers a short middle finger (**figure 14.4**). We can deduce the frequencies of the dominant and recessive alleles by observing the frequency of homozygous recessives, because this phenotype—short finger—reflects only one genotype. If 9 out of 100 individuals in a population have short fingers—genotype dd —the frequency is 9/100 or 0.09. Since dd equals q^2 , then q equals 0.3. Since $p + q = 1.0$, knowing that q is 0.3 tells us that p is 0.7.

Next, we can calculate the proportions of the three genotypes that arise when gametes combine at random:

$$\begin{aligned} \text{Homozygous dominant} &= DD \\ &= 0.7 \times 0.7 = 0.49 \\ &= 49 \text{ percent of individuals in generation 1} \end{aligned}$$

$$\begin{aligned} \text{Homozygous recessive} &= dd \\ &= 0.3 \times 0.3 = 0.09 \\ &= 9 \text{ percent of individuals in generation 1} \end{aligned}$$

$$\begin{aligned} \text{Heterozygous} &= Dd + dD \\ &= 2pq = (0.7)(0.3) + (0.3)(0.7) = 0.42 \\ &= 42 \text{ percent of individuals in generation 1} \end{aligned}$$

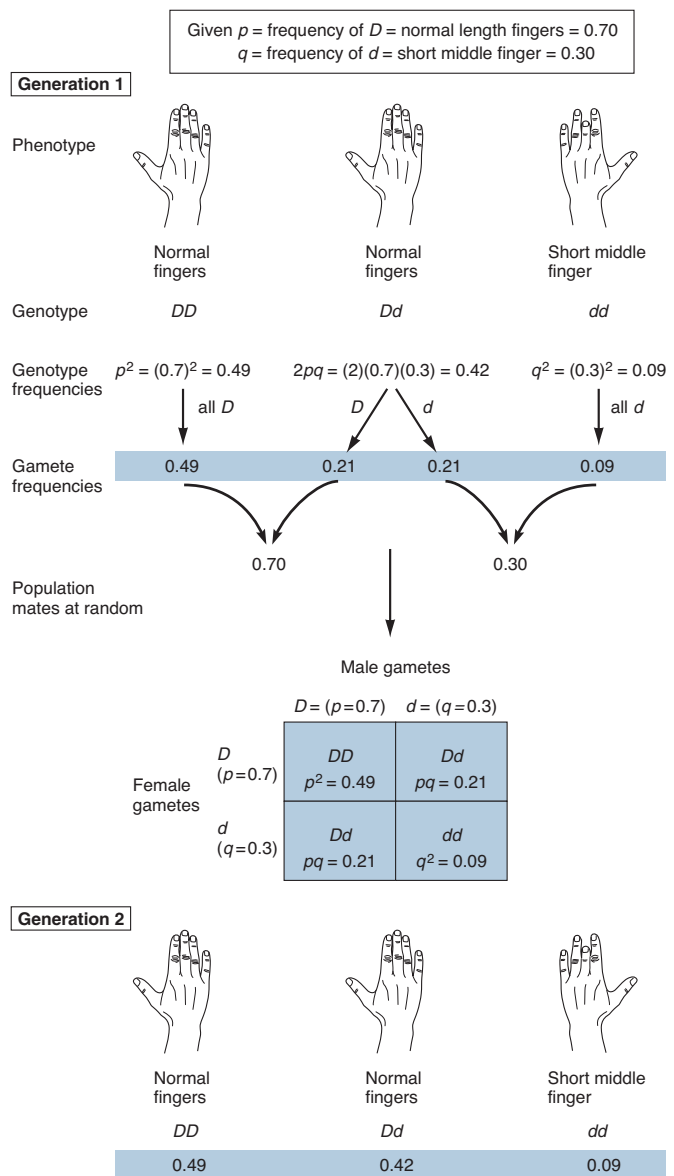


Figure 14.4 Hardy-Weinberg equilibrium. In Hardy-Weinberg equilibrium, allele frequencies remain constant from one generation to the next.

The proportion of homozygous individuals is calculated simply by multiplying the allele frequency for the recessive or dominant allele by itself. The heterozygous calculation is $2pq$ because there are two ways of combining a D with a d gamete—a D sperm with a d egg, and a d sperm with a D egg.

In this population, 9 percent of the individuals have a short middle finger. Now jump ahead a few generations, and assume that people choose mates irrespective of finger length. This means that each genotype of a female (DD , Dd , or dd) is equally likely to mate with each of the three types of males (DD , Dd , or dd), and vice versa. **Table 14.2** multiplies the genotype frequencies for each

Table 14.2 Hardy-Weinberg Equilibrium—When Allele Frequencies Stay Constant

POSSIBLE MATINGS			FREQUENCY OF OFFSPRING GENOTYPES		
Male	Female	Proportion in Population	<i>DD</i>	<i>Dd</i>	<i>dd</i>
0.49 <i>DD</i>	0.49 <i>DD</i>	0.2401 (<i>DD</i> × <i>DD</i>)	0.2401		
0.49 <i>DD</i>	0.42 <i>Dd</i>	0.2058 (<i>DD</i> × <i>Dd</i>)	0.1029	0.1029	
0.49 <i>DD</i>	0.09 <i>dd</i>	0.0441 (<i>DD</i> × <i>dd</i>)		0.0441	
0.42 <i>Dd</i>	0.49 <i>DD</i>	0.2058 (<i>Dd</i> × <i>DD</i>)	0.1029	0.1029	
0.42 <i>Dd</i>	0.42 <i>Dd</i>	0.1764 (<i>Dd</i> × <i>Dd</i>)	0.0441	0.0882	0.0441
0.42 <i>Dd</i>	0.09 <i>dd</i>	0.0378 (<i>Dd</i> × <i>dd</i>)		0.0189	0.0189
0.09 <i>dd</i>	0.49 <i>DD</i>	0.0441 (<i>dd</i> × <i>DD</i>)		0.0441	
0.09 <i>dd</i>	0.42 <i>Dd</i>	0.0378 (<i>dd</i> × <i>Dd</i>)		0.0189	0.0189
0.09 <i>dd</i>	0.09 <i>dd</i>	0.0081 (<i>dd</i> × <i>dd</i>)			0.0081
Resulting offspring frequencies:			0.49	0.42	0.09
			<i>DD</i>	<i>Dd</i>	<i>dd</i>

possible mating, which leads to offspring in the familiar proportions of 49 percent *DD*, 42 percent *Dd*, and 9 percent *dd*. This gene, therefore, is in Hardy-Weinberg equilibrium—the allele and genotype frequencies do not change from one generation to the next.

Key Concepts

1. For any two alleles of a gene in a population, the proportion of homozygous dominants equals the square of the frequency of the dominant allele (p^2), and the proportion of homozygous recessives equals the square of the frequency of the recessive allele (q^2). The proportion of heterozygotes equals $2pq$.
2. The frequency of the recessive allele equals the proportion of homozygous recessives plus one-half that of carriers, and the frequency of the dominant allele equals the proportion of homozygous dominants plus one-half that of carriers.
3. In Hardy-Weinberg equilibrium, allele frequencies remain constant from one generation to the next.

14.3 Applying Hardy-Weinberg Equilibrium

A young woman pregnant for the first time watches a television program about cystic fibrosis. Alarmed to learn about the daily treatments and possible complications, and that CF is the most common genetic disorder in her population group (Caucasian of European descent), the woman wonders what the risk is that

her child will have CF—even though there is no known history of the disorder in her or her partner’s families, which are from the same ethnic group. The Hardy-Weinberg equation can help to answer that question by determining the probability that the woman and her partner are carriers. If they are, then Mendel’s first law can be used to calculate the risk to offspring.

To derive carrier risks, the Hardy-Weinberg equation is applied to population statistics on genetic disease incidence. To determine allele frequencies for autosomal recessively inherited characteristics, we need to know the frequency of one genotype in the population. This is typically the homozygous recessive class, because its phenotype indicates its genotype.

The incidence (frequency) of an autosomal recessive disorder in a population is used to help calculate the risk that a particular person is a heterozygote. Returning to the example of CF, the incidence of the disease, and therefore also of carriers, may vary greatly in different populations (table 14.3).

Table 14.3 Carrier Frequency for Cystic Fibrosis

Population Group	Carrier Frequency
African Americans	1 in 66
Asian Americans	1 in 150
Caucasians of European descent	1 in 23
Hispanic Americans	1 in 46

CF affects 1 in 2,000 Caucasian newborns. Therefore, the homozygous recessive frequency— cc if c represents the disease-causing allele—is $1/2,000$, or 0.0005 in the population. This equals q^2 . The square root of q^2 is about 0.022 , which equals the frequency of the c allele. If q equals 0.022 , then p , or $1 - q$, equals 0.978 . Carrier frequency is equal to $2pq$, which equals $(2)(0.978)(0.022)$, or 0.043 —about 1 in 23. **Figure 14.5** summarizes these calculations.

Since there is no CF in the woman's family, her risk of having an affected child, based on population statistics, is low. The chance of *each* potential parent being a carrier is about 4.3 percent, or 1 in 23. The chance that *both* are carriers is $1/23$ multiplied by $1/23$ —or 1 in 529—because the probability that two independent events will occur equals the product of the probability that each event will happen alone. However, if they *are* both carriers, each of their children would face a 1 in 4 chance of inheriting the illness, based on Mendel's first law of gene segregation. Therefore, the risk that these two unrelated Caucasian individuals with no family history of CF will have an affected child is $1/4 \times 1/23 \times 1/23$, or 1 in 2,116.

For X-linked traits, different predictions of allele frequencies apply to males and females. For a female, who can be homozygous recessive, homozygous dominant, or a heterozygote, the standard Hardy-Weinberg equation of $p^2 \times 2pq \times q^2$ applies. However, in males, the allele frequency is the phenotypic frequency, because a male who inherits an X-linked recessive allele exhibits it in his phenotype.

The incidence of X-linked hemophilia A (see figure 6.8), for example, is 1 in 10,000 male (X^hY) births. Therefore, q (the frequency of the h allele) equals 0.0001 . Using the formula $p + q = 1$, the frequency of the wild type allele is 0.9999 . The incidence of carriers (X^HX^h), who are all female, equals $2pq$, or $(2)(0.0001)(0.9999)$, which equals 0.00019 ; this is 0.0002 , or 0.02 percent, which equals about 1 in 5,000. The incidence of a female having hemophilia A (X^hX^h) is q^2 , or $(0.0001)^2$, or about 1 in 100 million. **Figure 14.6** summarizes these calculations.

Neat allele frequencies such as 0.6 and 0.4 , or 0.7 and 0.3 , are unusual. In actuality, single-gene disorders are very rare, and so the q component of the Hardy-Weinberg equation contributes little. Because this means that the value of p approaches 1, the carrier frequency, $2pq$, is very close to $2q$. Thus, the

Cystic Fibrosis

incidence (autosomal recessive class) = $1/2,000 = 0.0005$

$$\therefore q^2 = 0.0005$$

$$\therefore q = \sqrt{0.0005} = 0.022$$

$$\therefore p = 1 - q = 1 - 0.022 = 0.978$$

$$\therefore \text{carrier frequency} = 2pq = (2)(0.978)(0.022) = 0.043 = 1/23$$

Figure 14.5 Calculating the carrier frequency given population incidence: Autosomal recessive.

Hemophilia A

incidence = $1/10,000$ male births = 0.0001

$$\therefore q = 0.0001$$

$$\therefore p = 1 - q = 1 - 0.0001 = 0.9999$$

$$\therefore \text{carrier frequency (females)} = 2pq = (2)(0.9999)(0.0001) = 0.00019 = \text{about } 1/5,000$$

$$\therefore \text{affected females} = q^2 = (0.0001)(0.0001) = 1/100 \text{ million}$$

Figure 14.6 Calculating the carrier frequency given population incidence: X-linked recessive.

carrier frequency is approximately twice the frequency of the rare, disease-causing allele.

Consider Tay-Sachs disease, which occurs in 1 in 3,600 Ashkenazim (Jewish people of eastern European descent). This means that q^2 equals $1/3,600$, or about 0.0003 . The square root, q , equals 0.017 . The frequency of the dominant allele (p) is then $1 - 0.017$, or 0.983 . What is the likelihood that an Ashkenazi carries Tay-Sachs disease? It is $2pq$, or $(2)(0.983)(0.017)$, or 0.033 . This is very close to double the frequency of the mutant allele (q), 0.017 . Modifications of the Hardy-Weinberg equation are used to analyze genes that have more than two alleles.

Key Concepts

1. Allele frequencies in populations can be inferred from the frequency of homozygous recessive individuals (q^2). The values of q and p can then be deduced and the Hardy-Weinberg equation applied to predict the frequency of carriers.
2. For X-linked traits, the frequency of the recessive phenotype in males is q , and in females q^2 .
3. For very rare inherited disorders, p approaches 1, so the carrier frequency is approximately twice the frequency of the disease-causing allele ($2q$).

14.4 DNA Profiling Uses Hardy-Weinberg Assumptions

Hardy-Weinberg equilibrium is useful and interesting in a theoretical sense to understand the conditions necessary for evolution to occur. These calculations are also the foundation of DNA profiling. Use of Hardy-Weinberg equilibrium in DNA profiling is based on the fact that the equation applies to parts of the genome that do not affect the phenotype, and are therefore not subject to natural selection. Short repeated sequences that are not part of a protein-encoding gene fall into this category. Variability in such sequences can be used to identify

individuals if the frequencies are known in particular populations.

Recall from chapter 12 that repeated sequences are scattered throughout the genome. Copy number variants (the number of copies of a particular repeat) can be followed, as alleles, to identify an individual. The person is classified as a heterozygote or a homozygote based on the number of copies of the same repeat at the same chromosomal locus on the two homologs. A homozygote has the same number of repeats on both homologs, such as individual 2 in **figure 14.7**. A heterozygote has two different repeat sizes, such as the other two individuals in the figure. The copy numbers are distributed in the next generation according to Mendel’s law of segregation. A child of individual 1 and individual 2 in figure 14.7, for example, could have any of the two possible combinations of the parental copy numbers, one from each parent: 2 and 3, or 4 and 3.

DNA profiling was pioneered on detecting copy number variants of very short repeats and using them to identify or distinguish individuals. In general, the technique calculates the probability that certain combinations of repeat numbers will be in two DNA sources by chance. For example, if a DNA profile of skin cells taken from under the fingernails of an assault victim matches the profile from a suspect’s hair, and the likelihood is very low that those two samples would match by chance, that is strong evidence of guilt rather than a coincidental similarity. DNA evidence is more often valuable in excluding a suspect, and should be considered along with other types of evidence.

Although obtaining a DNA profile is a molecular technique, interpreting it requires statistical analysis of population data. Two types of repeats are used in forensics and in identifying victims of disasters: **variable number of tandem repeats** (VNTRs), and **short tandem repeats** (STRs). **Table 14.4** compares them.

DNA Profiling Began with Forensics

Sir Alec Jeffreys at Leicester University in the United Kingdom invented DNA profiling (then called DNA fingerprinting) in the 1980s. He detected differences in numbers of VNTRs among individuals by cutting DNA with restriction enzymes. These

Type	Repeat Length	Distribution	Example	Fragment Sizes
VNTRs (minisatellites)	10–80 bases	not uniform	TTCGGGTTG	50–1,500 bases
STRs (microsatellites)	2–10 bases	more uniform	ACTT	50–500 bases

enzymes naturally protect bacteria by cutting foreign DNA, such as DNA from viruses, at specific short sequences. They are used as “molecular scissors” in biotechnology, as discussed in chapter 19. Jeffreys measured DNA fragments using a technique called agarose gel electrophoresis, described in **Reading 14.1**. The different-sized fragments that result from “digesting” DNA with these enzymes are called restriction fragment length polymorphisms (RFLPs, pronounced “riflips”).

In the technique that Jeffreys used, DNA pieces migrate through a jellylike material (agarose or the more discriminating polyacrylamide) when an electrical field is applied. A positive electrode is placed at one end of the gel strip, and a negative electrode at the other. The DNA pieces, carrying negative charges because of their phosphate groups, move toward the positive pole. The pieces migrate according to size, with the shorter pieces moving faster and thus traveling farther in a given time. The pattern that forms when the different-sized fragments stop moving, with the shorter fragments closer to the positive pole and the longer ones farther away, creates a distinctive DNA pattern, or profile, that looks like a strip of black smears. An individual who is heterozygous for a repeat copy number variant will have two bands for that locus, as shown in **figure 14.8** for Individuals 1 and 3 from figure 14.7. A locus for which an individual is homozygous has only one corresponding band (Individual 2), because both DNA pieces are the same size.

Jeffreys’ first cases proved that a boy was the son of a British citizen so that he could enter the country, and freed a man jailed for raping two schoolgirls. Then in 1988, Jeffreys’ approach matched DNA profiles from suspect Tommie Lee Andrews’ blood cells to sperm cells left on his victim in a notorious rape case. Jeffreys also used DNA profiling to demonstrate that Dolly, the Scottish sheep, was truly a clone of the 6-year-old ewe that donated her nucleus (**figure 14.9**).

DNA can be obtained from any cell with a nucleus. Common sources include cells in hair, blood, skin, secretions, or the inside of the cheek. DNA sequences other than VNTRs are used when sample DNA is scarce. STRs are used when DNA is fragmented, such as in evidence from terrorist attacks and natural disasters. Their smaller size makes them more likely to persist in degraded DNA. STRs are amplified using the polymerase chain reaction (see chapter 19).

If DNA is extremely damaged, such that even STRs are obliterated,



Figure 14.7 DNA profiling detects differing numbers of repeats at specific chromosomal loci. Individuals 1 and 3 are heterozygotes for the number of copies of a 5-base sequence at a particular chromosomal locus. Individual 2 is a homozygote, with the same number of repeats on the two copies of the chromosome. (Repeat number is considered an allele.)

Reading 14.1

DNA Profiling: Molecular Genetics Meets Population Genetics

DNA profiling is a standard and powerful tool in forensic investigations, agriculture, paternity testing, and historical investigations. Until 1986, it was unheard of outside of scientific circles. A dramatic rape case changed that.

Tommie Lee Andrews watched his victims months before he attacked so that he knew when they would be home alone. On a balmy Sunday night in May 1986, Andrews awaited Nancy Hodge, a young computer operator at Disney World in Orlando, Florida. The burly man

surprised her when she was in her bathroom removing her contact lenses. He covered her face, then raped and brutalized her repeatedly.

Andrews was very careful not to leave fingerprints, threads, hairs, or any other indication that he had ever been in Hodge's home. But he left DNA. Thanks to a clearthinking crime victim and scientifically savvy lawyers, Andrews was soon at the center of a trial that would judge the technology that helped to convict him.

After the attack, Hodge went to the hospital, where she provided a vaginal secretion sample containing sperm. Two district attorneys who had read about DNA testing sent some of the sperm to a biotechnology company that extracted DNA and cut it with restriction enzymes. The sperm's DNA pieces were then mixed with labeled DNA probes that bound to complementary sequences.

The same extracting, cutting, and probing of DNA was done on white blood cells from Hodge and Andrews, who had been held as a suspect in several assaults. When the radioactive DNA pieces from each sample, which were the sequences where the probes had bound, were separated and displayed by size, the resulting pattern of bands—the DNA profile—matched exactly for the sperm sample and Andrews' blood, differing from Hodge's DNA (**figure 1**).

Andrews' allele frequencies were compared to those for a representative African American population. At his first trial in November 1987, the judge, perhaps fearful that too much technical information would overwhelm the jury, did not allow the prosecution to cite population-based statistics. Without the appropriate allele frequencies, DNA profiling was just a comparison of smeary lines on test papers to see whether the patterns of DNA pieces in the forensic sperm sample looked like those for Andrews' white blood cells. Although population-based statistics indicated that the possibility that Andrews' DNA would match the evidence by chance was 1 in 10 billion, the prosecution could not mention this. After a mistrial was declared, the prosecution cited the precedent of using population statistics to derive databases on standard blood types. So when Andrews stood trial just 3 months later for raping a different woman, the judge permitted population analysis. Andrews was convicted.

Although DNA profiling is widely held to be an extremely accurate method to match a suspect to a sample, setbacks still happen. In June 2009, the Supreme Court ruled against prisoner William Osborne's repeated requests for further DNA testing, in a case very similar to the one described in the chapter opener. The crime was committed in Alaska, one of a very few states not to allow DNA testing of prisoners. In these states, prisoners must take their cases to the national level. However, Peter Neufeld, co-director of the Innocence Project, said that the ruling would have limited impact on the availability of DNA profiling because most states, and many prosecutors, allow requested DNA testing.

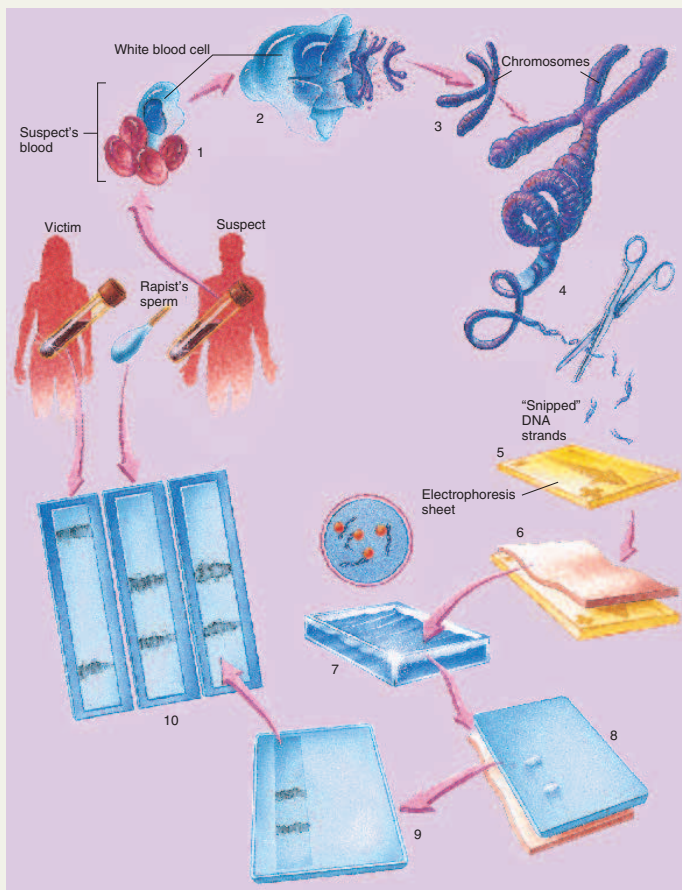


Figure 1 DNA profiling. A blood sample (1) is collected from the suspect. White blood cells are separated and burst open (2), releasing DNA (3). Restriction enzymes snip the strands into fragments (4), and electrophoresis aligns them by size in a groove on a sheet of gel (5). The resulting pattern of DNA fragments is transferred to a nylon sheet (6). It is then exposed to radioactively tagged probes (7) that bind the DNA areas used to establish identity. When the nylon sheet is placed against a piece of X-ray film (8) and processed, black bands appear where the probes bound (9). This pattern of bands is a DNA profile (10). It may be compared to the victim's DNA pattern, the rapist's DNA obtained from sperm cells, and other biological evidence. Today fluorescent labels are used.

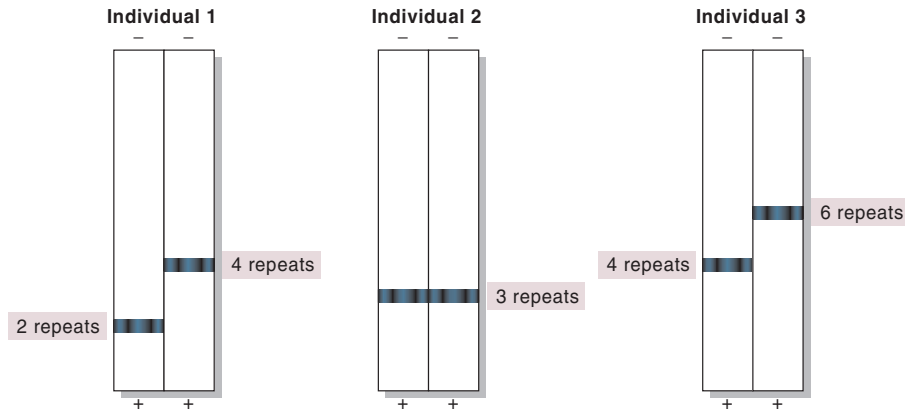


Figure 14.8 DNA profiles. DNA fragments that include differing numbers of copies of the same repeat migrate at different speeds and stop moving at different points on a strip of polyacrylamide gel. These gels correspond to the individuals represented in figure 14.7. Actual DNA profiles typically scan up to 25 repeats on different chromosomes.

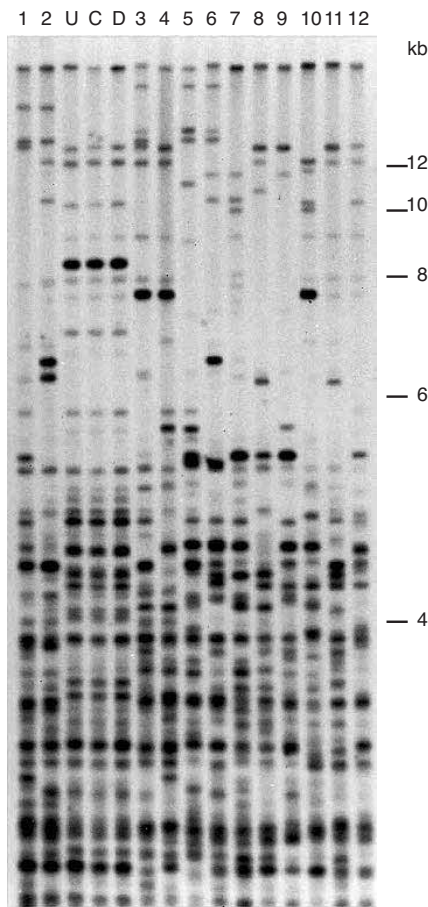


Figure 14.9 Comparing DNA profiles. These DNA profiles compare the DNA of Dolly the cloned sheep (lane D), fresh donor udder tissue (U), and cultured donor udder tissue (C). The other twelve lanes represent other sheep. The match between Dolly and the two versions of her nucleus donor is obvious.

mitochondrial DNA (mtDNA) is often used instead, particularly two regions of repeats that are highly variable in populations. Because a single cell can yield hundreds or thousands of copies of the mitochondrial genome, even vanishingly small forensic samples can yield this DNA.

MtDNA analysis was critical in analyzing evidence from the September 11 terrorist attacks, most of which was extremely degraded. A more bizarre application was the case of the “voodoo child.” The evidence was a boy’s torso found floating in the River Thames in east London. The name reflects the contents of the stomach, which suggested he had been the victim of a ritualistic killing. When the DNA profile of nuclear DNA from the torso did not match that of missing English children, investigators widened the search by using a global mtDNA database. This search led to the boy’s homeland, southwestern Nigeria. He had been kidnapped, enslaved, and beheaded. Several suspects were arrested, thanks to tracking the torso to Africa.

Commercially available software enables researchers to integrate different types of DNA profiling data. For forensic applications, the FBI’s Combined DNA Index System (CODIS) shares DNA profiles electronically among local, state, and federal crime laboratories. More than 3 million DNA profiles are stored, and searching CODIS for DNA profiles has led to more than 22,000 “cold hits”—identifying a suspect from DNA alone. CODIS uses the thirteen STRs shown in **figure 14.10**. The probability that any two individuals have the same thirteen STR markers by chance is 1 in 250 trillion. Therefore, identity at all thirteen sites is a virtual match, but just one mismatch disproves identity.

Using Population Statistics to Interpret DNA Profiles

In forensics in general, the more clues, the better. Therefore, the power of DNA profiling is greatly expanded by tracking repeats on several chromosomes. The numbers of copies of a repeat are assigned probabilities (likelihood of being

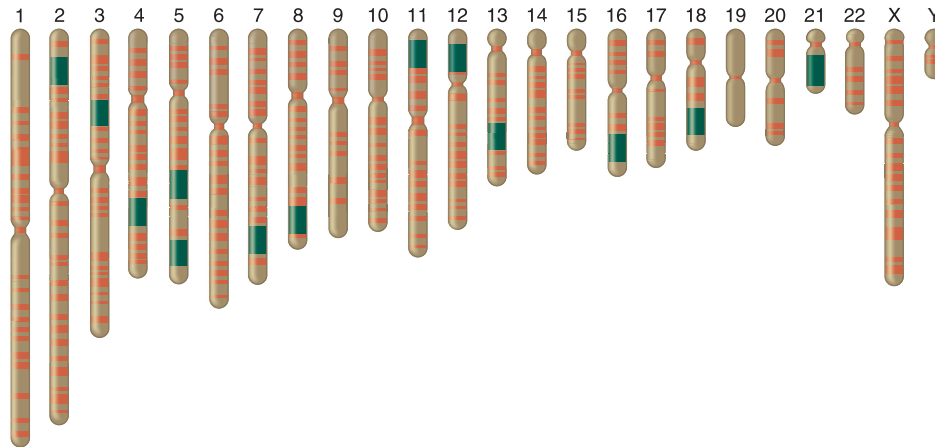


Figure 14.10 DNA profiling. A minimum of thirteen sites in the genome are compared to rule out suspects in crimes. The green bands indicate the thirteen original CODIS sites. More are increasingly being used.

present) based on their observed frequencies in a particular population. Considering repeats on different chromosomes makes it possible to use the product rule to calculate the probabilities of particular combinations of repeat numbers occurring in a population, based on Mendel's law of independent assortment.

The Hardy-Weinberg equation and the product rule are used to derive the statistics that back up a DNA profile. First, the pattern of fragments indicates whether an individual is a homozygote or a heterozygote for each repeat, because a homozygote only has one band representing that locus. Genotype frequencies are then calculated using parts of the Hardy-Weinberg equation. That is, p^2 and q^2 denote each of the two homozygotes for a two-allele repeat, and $2pq$ represents the heterozygote. Then the frequencies are multiplied.

Table 14.5 shows an example of multiplying frequencies of different repeat numbers. The result is the probability that this particular combination of repeat sizes would occur in a particular population. Logic then enters the equation. If the combination is very rare in the population the suspect comes from, and if it is found both in the suspect's DNA and in crime scene evidence, such as a rape victim's body or the stolen property in table 14.5, the suspect's guilt appears highly likely. **Figure 14.11** summarizes the procedure.

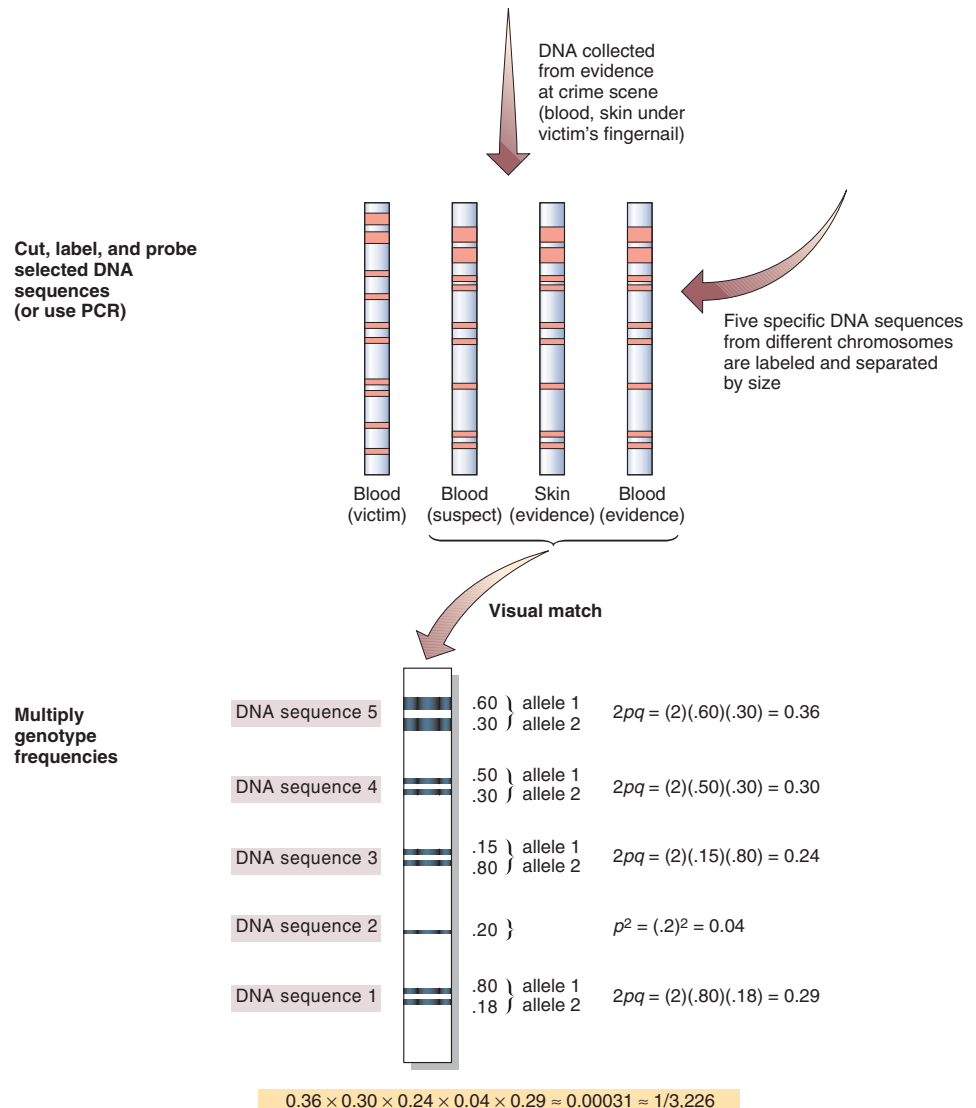
For the sequences used in DNA profiling, Hardy-Weinberg equilibrium is assumed. When it doesn't apply, problems can arise. For example, the requirement of non-random mating for Hardy-Weinberg equilibrium wouldn't be met in a community with a few very large families where

Table 14.5 Multiplied Frequencies of Different Repeat Numbers

The Case: A famous painting has been stolen from a gallery. The thief planned the crime carefully, but as she was removing the painting from its display, she sneezed. She averted her face, but a few tiny droplets hit the wall. Detectives obtained a DNA profile using six repeat alleles, from different chromosomes, for DNA in nose lining cells in the droplets. Then they compared the profile to those compiled for eight people in the vicinity, all women, who had been identified by hidden camera. (Assume the suspects are in the same ethnic group.) Most of the samples matched at two to four sites, but one matched at all six. She was the crook. Notice how the probability of guilt increases with the number of matches. Matching for the very rare allele #3 is particularly telling.

Allele	Repeat	Frequency	Cumulative Multiplied Frequencies
1	ACT on chromosome 4	1/60	
2	GGC on chromosome 17	1/24	$1/60 \times 1/24 = 1/1,440$
3	AAGCTA on chromosome 14	1/1,200	$1/1,440 \times 1/1,200 = 1/1,728,000$
4	GGTCTA on chromosome 6	1/11	$1/1,728,000 \times 1/11 = 1/19,008,000$
5	ATACGAGG on chromosome 9	1/40	$1/19,008,000 \times 1/40 = 1/760,320,000$
6	GTA on chromosome 5	1/310	$1/760,320,000 \times 1/310 = 1/235,699,200,000$

Figure 14.11 To solve a crime. A man was found brutally murdered, with bits of skin and blood beneath a fingernail. The bits were sent to a forensics lab as evidence, where the patterns of five DNA sequences were compared to patterns in blood from the victim as well as blood from a man being held as a suspect. The pattern for the crime scene evidence matched that for the suspect visually, but that wasn't sufficient. Allele frequencies from the man's ethnic group were used in the Hardy-Weinberg equation, yielding the probability that his DNA matched that of the skin and blood under the murdered man's fingernail by chance.



Conclusion: The probability that another person in the suspect's population group has the same pattern of these alleles is approximately 1 in 3,226.

distant relatives might inadvertently marry each other—a situation in many small towns. A particular DNA profile for one person might be shared by his or her cousins. In one case, a young man was convicted of rape based on a DNA profile—which he shared with his father, the actual rapist. Considering a larger number of repeat sites can guard against such complications. If more repeat sites had been considered in the rape case, chances are that they would have revealed a polymorphism that the son had inherited from his mother, but that the father lacked. This would have indicated that the son was not guilty, but a close male relative might be.

The accuracy and meaning of a DNA profile depend upon the population that is the source for the allele frequencies. If populations are too broadly defined, then allele frequencies are typically low, leading to very large estimates of the likelihood that a suspect matches evidence based on chance. In one oft-quoted trial, the prosecutor concluded, *The chance of the DNA fingerprint of the cells in the evidence matching blood of the defendant by chance is 1 in 738 trillion*. The numbers were accurate, but some population geneticists questioned the validity of the databases. Did they really reflect the gene pool compositions of actual populations? By 1991, several judges had rejected DNA evidence because population geneticists had testified that the databases greatly oversimplify human population structure. Therefore, the odds that crime scene DNA matched suspect DNA were not as reliable as originally suggested.

The first DNA profiling databases neatly shoehorned many different groups into just three—Caucasian, black, or Hispanic—designations not necessarily biologically meaningful. People from Poland, Greece, or Sweden would all be considered white, and a dark-skinned person from Jamaica and one from Somalia would be lumped together as blacks. Perhaps the most incongruous of all were the Hispanics. Cubans and Puerto Ricans are part African, whereas people from Mexico and Guatemala have mostly Native American gene variants. Spanish and Argentinians have neither black African nor Native American genetic backgrounds. Yet these diverse peoples were considered a single population! Other groups were left out, such as Native Americans and Asians. Ultimately, analysis of these three databases revealed significantly more homozygous recessives for certain polymorphic genes than the Hardy-Weinberg equation would predict, confirming what many geneticists had suspected—allele frequencies were not in equilibrium.

Giving meaning to the allele frequencies necessary to interpret DNA profiles requires more restrictive ethnic databases. A frequency of 1 in 1,000 for a particular allele in all whites may actually be much higher or lower in, for example, only Italians, because they (and many others) tend to marry among themselves. On the other hand, narrowly defined ethnic databases may be insufficient to interpret DNA profiles from people of mixed heritages, such as someone whose mother was Scottish/French and whose father was Greek/German.

We may need to develop mathematical models to account for real, mixed population structures. Perhaps the first step will be to understand the forces that generate genetic substructures within more broadly defined populations, which means taking into account history and human nature. Chapter 15 explores these factors.

Using DNA Profiling to Identify Victims

Early editions of this textbook described using DNA profiling to identify human remains from plane crashes—at first the largest application of the technology. Then terrorist attacks and natural disasters took the scope of DNA profiling to a new level.

Many cases that use DNA profiling help each other. Techniques perfected in the case of murdered 6-year-old beauty queen JonBenet Ramsey, for example, were used on some of the 13,000 bone splinters in the wake of the 9/11/01 terrorist attacks in New York City. Further improvements made in New York City were applied to identifying bodies in mass graves in Peru years later. The world, it seems, has no shortage of disasters.

Identifying World Trade Center Victims

In late September, 2001, Myriad Genetics, a company in Salt Lake City that normally provides breast cancer tests, received three unusual types of DNA samples:

- evidence from the World Trade Center
- cheek brush scrapings from relatives of people missing from the site
- “reference samples” from the victims’ toothbrushes, razors, and hairbrushes.

Technologists analyzed the DNA for copy numbers of the thirteen standard STRs as well as the sex chromosomes. If the STR pattern of crime scene evidence matched DNA from a victim’s toothbrush, identification was fairly certain. Myriad sent its results to the New York State Forensic Laboratory, where investigators matched family members to victims.

STR analysis worked on pieces of soft tissue, but bone bits that persisted despite the ongoing fire at the site required harder mtDNA analysis. Although 20,000 tissue pieces were profiled, nearly half of the victims have not been identified. However, new evidence is found now and then.

DNA profiling provides much more reliable information on identity than traditional forensic identifiers such as dental patterns, scars, and fingerprints, and clues such as jewelry, wallets, and rolls of film found with the victim. Consider the case of Jose Guadalupe and Christopher Santora, two of fifteen firefighters lost from one engine company on September 11, 2001. Rescue workers brought a body found beside a firetruck next to the destroyed towers to the Medical Examiner’s office on September 13th. Other firefighters identified the remains as belonging to Guadalupe based on where the body had been found, because he had been the driver of the firetruck. The body also had a gold chain that the men recognized, and X rays

revealed a birth defect in the neck bone that he was known to have had. Guadalupe was buried on October 1—but it wasn't Guadalupe in the grave. Santora had the same necklace and the same neck condition! A DNA sample taken from the buried man's remains and from Santora's relatives matched.

Identifying Natural Disaster Victims

Different types of disasters present different challenges for DNA profiling (table 14.6 and figure 14.12). The people caught in the September 11 terrorist attacks died by fire; those caught in the Indian Ocean tsunami of 2004 or in hurricane Katrina in 2005 died by water. Whereas New York City workers searched rubble for remains, the 250,000-plus bodies strewn about by the tsunami were everywhere. Rather than hunting for tissue bits out in the open, tsunami workers had to exhume bodies that had been buried in haste to stem the spread of infectious disease. Those remains that were accessible after the waves hit quickly decayed in the hot, wet climate. These conditions, combined with the lack of roads and labs, led to 75 percent of the bodies being identified by standard dental record analysis, and 10 percent from fingerprints. Fewer than half of 1 percent of the victims were identified by their DNA.

Forensic scientists had learned from 9/11 the importance of matching victim DNA to that of relatives, to avoid errors when two people matched at several genome sites by chance. In New York City, many of those relatives were from nearby neighborhoods; in the Asian disaster, 12 countries were directly affected and victims came from 30 countries. Entire families were washed away, leaving few and many times no relatives to provide DNA, even if everyday evidence such as toothbrushes had remained. Conditions in New Orleans were similar, but on a smaller scale. The Pakistan earthquake of 2005 combined the challenges of the 9/11 attacks, the tsunami, and the hurricane: collecting samples for DNA profiling was very difficult.

To compensate for the barriers to implementing DNA profiling in mass disasters, Sir Alec Jeffreys advised

Table 14.6

Challenges to DNA Profiling in Mass Disasters

- Climate that hastens decay
- Inability to reach remains
- No laboratory facilities
- Number of casualties
- Lack of relatives
- Destruction of personal item evidence
- Poor DNA quality (too fragmented, scarce, degraded)
- Lack of availability of DNA probes and statistics for population



Figure 14.12 Challenges to DNA profiling. On a small scale, such as a murder investigation, DNA profiling is enormously valuable. For large-scale disasters, many practical constraints limit the utility of the technique. The tsunami in southeast Asia in late 2004, shown here, washed away nearly all evidence; most of what little remained degraded in the heat. As a result, traditional forensic techniques, such as comparing dental records and fingerprints, proved more useful.

assessing 15 to 20 repeat (copy number variant) sites, rather than the usual 13, and some investigators recommend upping the number to 50. Tragic as these disasters were, they have spurred forensic scientists to develop ways to better integrate many types of evidence, including that found in DNA sequences.

Reuniting Holocaust Survivors

A happier use of DNA profiling is to reunite families who were torn apart in the Holocaust of World War II. The DNA Shoah project has established a DNA database of many of the 300,000 or so survivors, including some of the 10,000-plus Holocaust orphans. (*Shoah* is Hebrew for *holocaust*.) Michael Hammer and his colleagues at the University of Arizona compare the data to DNA profiles from human remains unearthed in various building projects in parts of Europe where the mass killings occurred. The challenges in reuniting Holocaust families combine those of the 9/11 and tsunami investigations: degraded DNA and few surviving relatives and descendants. The Shoah project is linking the past to the present by matching DNA profiles.

Identifying an Individual's DNA in Mixtures

Natural disasters and crime scenes can leave investigators with DNA from many sources. In many situations, DNA from different individuals contributes different proportions to the evidence. Researchers applied principles of population genetics,

along with powerful algorithms, to develop a way to identify an individual's DNA in a mixture. The approach uses a microarray to probe DNA fragments for many thousands of SNPs in three sources of DNA:

- DNA from a “person of interest,” such as from a toothbrush of a disaster victim
- DNA from the mixture (forensic sample)
- DNA from a “reference population” that matches the person of interest, taking into account ancestry, including mixed ancestry such as northern European with sub-saharan African.

Genotypes are determined for the two copies of every SNP (from the two copies of the chromosome in a presumably diploid cell). Places in the genome where the mixture DNA and reference population DNA SNPs differ are noted, as well as which SNPs the person of interest's DNA matches. Overall, if the person's DNA tracks significantly more with the mixture DNA than with his or her reference DNA, then the DNA is likely, indeed, in the mixture. The power of the approach comes from the large number of SNPs, which provide points in the genome for comparison. In simulations as well as experiments, researchers found that using up to 50,000 SNPs can identify a person's DNA that contributes as little as 0.1 percent to a mixture!

The ability to detect an individual's DNA in a mixture brought up a huge, unexpected bioethical concern: If this could be done for forensic samples, then people who participate in genome-wide association studies can also be identified. (This was discussed in *Bioethics: Choices for the Future* in chapter 1.) When researchers at the National Institutes of Health read the study about identifying DNA in mixtures, they immediately limited Internet access to genome-wide association databases, to protect the privacy of participants—the next topic.

Genetic Privacy

Before the information age, population genetics was an academic discipline that was more theoretical than practical. Today, with the combination of information technology, genome-wide association studies, genome sequencing, and shortcuts to identify people by SNP or copy number patterns, population genetics presents a powerful way to identify individuals. This new view of population genetics raises both personal and societal challenges (see *Bioethics: Choices for the Future* on page 279).

The human genome is 3.2 billion bits of information, each of which can be one of four possibilities—that's a huge capacity for diversity. Our genomes can vary many more ways than there are people—about 10 billion worldwide. Given these daunting numbers, one only need consider 30 to 80 genome sites to uniquely describe each person. This is why forensic tests can compare only 10 to 15 or so loci (sites in the genome) to rule out or establish identity.

The ease of assigning highly individualized genetic nametags may be helpful in forensics, but it poses privacy issues. Consider a “DNA dragnet,” a forensic approach of taking DNA profiles of all residents of a town where a violent crime is unsolved. Sir Alec Jeffreys in the U.K. conducted some of the first DNA dragnets in the late 1980s. The largest to date occurred in 1998 in Germany, where more than 16,000 men had their DNA profiled in a search for the man who raped and murdered an 11-year-old. The dragnet indeed caught the killer.

A more recent DNA dragnet happened in the small town of Truro, near the tip of Cape Cod, Massachusetts. Writer Christa Worthington was brutally murdered in January 2002, a knife driven completely through her heart into the floorboards, and her toddler daughter found at her side, trying to mop up the blood. Only 790 men lived in the seaside village in the winter. DNA from semen in her body did not match samples in any criminal databases. Three years later, on the advice of federal authorities, police began asking men at Truro's few winter gathering places to provide cheek swabs for DNA testing. There was no requirement to do so, but a record was being kept of all who refused, and everyone knew it. Several citizens filed complaints with the American Civil Liberties Union, but most of Truro's male residents complied—including the trash collector who was convicted of the crime in 2006.

The Hardy-Weinberg equilibrium that makes DNA profiling possible is extremely rare in the real world, for most genes. The next chapter considers the familiar circumstances that change allele frequencies.

Key Concepts

1. DNA profiles are based on copy number variants.
2. Population statistics are applied to determine the probability that the same pattern would occur by chance in two individuals.
3. A limitation of the method is that databases may not adequately represent real populations. Developing narrower ethnic databases and considering historical and social factors may make population statistics more realistic.
4. DNA profiling of nuclear and mitochondrial DNA was performed on evidence from the September 11, 2001 terrorist attacks and the 2004 tsunami, and used in reuniting relatives separated in the Holocaust.
5. Identifying DNA in a mixed sample is possible by comparing many SNPs in the forensic sample to those in the relevant reference populations, and detecting overrepresentation.
6. Each person has a unique genetic signature (except multiples).
7. DNA profiling introduces privacy issues.

Bioethics: Choices for the Future

Population Biobanks

More than a dozen nations are recording and scrutinizing genetic, genealogical, lifestyle, and health information on their citizens to discover and archive the inherited and environmental influences on common disorders. These “biobank” projects vary in how people participate, but they raise similar concerns: Who will have access to the information? How can people benefit from providing it? How might it be abused?

Ideally, a biobank must meet several criteria. It should

- have data and tissue samples from at least 500,000 people.
- draw conclusions based on a population that is representative of the nation.
- have clinical information collected over many years.
- include family trees that link generations.
- compare results to those of other populations to validate DNA-disease associations.

The first country to systematically collect genetic information in a population was Iceland, where many citizens can trace their families back more than a thousand years. In 1998, a company called deCODE Genetics received government permission to collect existing health and genealogy records and to add DNA sequence data. Participation in the database is presumed—citizens must “opt out” of the project. Despite initial concerns (mostly from outside Iceland) that the populace would feel pressured to participate, that hasn’t been the case. The database includes 95 percent of everyone who has lived in the nation since 1703, when the first census was conducted. DeCODE has used the information to identify genes that contribute to more than 25 common disorders by identifying parts of the genome that people with these disorders uniquely share, then finding genes in these regions whose functions could explain the symptoms.

Some biobanks sample a population. CARTaGENE is randomly sampling 1 percent of Quebec’s citizens. Researchers in the United Kingdom are recruiting 500,000 individuals between the ages of 45 and 69, when many common illnesses begin, searching for connections among DNA sequence variants, health, and lifestyle characteristics as the population ages.

The United States is planning a similar project that will follow 500,000 citizens to assess genetic and environmental influences on common health conditions. The effort is controversial because 120,000 children will be included. So far, townhall meetings to introduce the plans have evoked concern about whether children would benefit from participating or be stressed, and how informed consent would be handled over time.

Some projects focus on specific medical conditions. The Estonian Genome Foundation uses registries for patients with cancer, Parkinson disease, diabetes mellitus, or osteoporosis. When patients show up for appointments, they learn about the project and are asked for details of their health histories and to donate DNA. Researchers then match variations in the DNA sequence to particular medical conditions. The United States has a similar program called GAIN (Genetic Association Information Network) that has DNA profiles (SNPs and copy number variants) from tissue sampled from 1,000 to 2,000 patients who have one of six common disorders. It is comparing these profiles to DNA profiles from the same number of healthy volunteers. The six conditions are psoriasis, schizophrenia, bipolar disorder, depression, diabetes, and attention deficit hyperactivity disorder. They were chosen based on the “highest likelihood of success” in identifying new drug targets.

Questions for Discussion

1. What do you think the minimal age should be for participation in a biobank?
2. How should privacy of participants be maintained?
3. Do you think that participation should be mandatory, as it is in Iceland? What are the risks and benefits of requiring participation?
4. If DNA testing reveals that an individual has inherited a mutation that could affect health, should blood relatives who could also have the mutation be notified?
5. Can you foresee any dangers of a biobank that uses genetic information? How can a problem be averted?

The First Biobanks

Biobank	Population	Website
CARTaGENE	Canada	http://www.cartagene.qc.ca/
DeCODE Genetics	Iceland	http://www.decode.com/
Estonian Genome Project	Estonia	http://www.geenivaramu.ee/index.php=show=main&lang=engl
GenomEUtwin	Europe	http://www.genomeutwin.org
UK Biobank	UK	http://www.ukbiobank.ac.uk/
The Genographic Project	Anyone	https://www3.nationalgeographic.com/genographic/index.html
BioBank Japan Project	Japanese	http://www.src.riken.jp/english/project/person/

Summary

14.1 Population Genetics Underlies Evolution

1. A **population** is a group of interbreeding members of the same species in a particular area. Their genes constitute the **gene pool**.
2. Population genetics considers allele, genotype, and phenotype frequencies to reveal microevolution. Phenotypic frequencies can be determined empirically, then used in algebraic expressions to derive other frequencies.
3. Genotype frequencies change if migration, nonrandom mating, genetic drift, mutations, or natural selection operate. In **Hardy-Weinberg equilibrium**, frequencies are not changing.

14.2 Constant Allele Frequencies

4. Hardy and Weinberg proposed an algebraic equation to explain the constancy of allele frequencies. This would show why dominant traits do not increase and recessive traits do not decrease in populations. The Hardy-Weinberg equation is a binomial expansion used to represent genotypes in a population.
5. Hardy-Weinberg equilibrium is demonstrated by following gamete frequencies as they recombine in the next generation. In equilibrium, these genotypes remain constant if evolution is not occurring. When the equation $p^2 + 2pq + q^2$ represents a gene with one dominant and one recessive allele, p^2 corresponds to the frequency of homozygous

dominant individuals; $2pq$ stands for heterozygotes; and q^2 represents the frequency of the homozygous recessive class. The frequency of the dominant allele is p , and of the recessive allele, q .

14.3 Applying Hardy-Weinberg Equilibrium

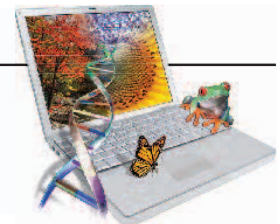
6. If we know either p or q , we can calculate genotype frequencies, such as carrier risks. Often such information comes from knowing the q^2 class, which corresponds to the frequency of homozygous recessive individuals in a population.
7. For X-linked recessive traits, the mutant allele frequency for males equals the trait frequency. For very rare disorders or traits, the value of p approaches 1, so the carrier frequency ($2pq$) is approximately twice the frequency of the rare trait (q).

14.4 DNA Profiling Uses Hardy-Weinberg Assumptions

8. Repeats (**VNTRs** and **STRs**) that do not encode protein are presumably in Hardy-Weinberg equilibrium and can be compared to establish individual DNA profiles.
9. To obtain a **DNA profile**, determine repeat numbers (using RFLPs or PCR) and multiply population-based allele frequencies to derive the probability that profiles from two sources match by chance.
10. People vary genetically in more ways than there are people.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

1. What is a population? List three populations.
2. "We like him, he seems to have a terrific gene pool," say the parents upon meeting their daughter's boyfriend. Why doesn't their statement make sense?
3. Explain the differences among an allele frequency, a phenotypic frequency, and a genotypic frequency.
4. What does Hardy-Weinberg equilibrium mean?
5. Why is Hardy-Weinberg equilibrium more a theoretical state than a common, real situation for genes that affect the phenotype?
6. What are the conditions under which Hardy-Weinberg equilibrium cannot be met?
7. Why is knowing the incidence of a homozygous recessive condition in a population important in deriving allele frequencies?
8. For a forensics case, why would tracking VNTR sequences provide a more reliable identification than tracking STRs?
9. Why are specific population databases needed to interpret DNA profiles?
10. How is the Hardy-Weinberg equation used to predict the recurrence of X-linked recessive traits?
11. What is the basis of assigning a probability value to a particular copy number variant?
12. Under what circumstances is analysis of repeats in mtDNA valuable?
13. Describe the following ways to identify or distinguish among individuals at the DNA level:
 - a. VNTRs
 - b. STRs
 - c. SNPs (see chapters 7 and 12)

- d. mtDNA
e. RFLPs
14. Why do forensic scientists using DNA profiling have to be extra careful in their analyses when the victim and suspect are blood relatives?
15. How did the challenges differ for profiling DNA in evidence from the September 11, 2001 attacks, the 2004 tsunami, and in matching Holocaust victims to survivors?
16. Suggest a novel use of DNA profiling.

Applied Questions

- Give three examples of how population genetics can be observed in everyday life.
- Why are Hardy-Weinberg calculations more complicated if a gene has many alleles that affect the phenotype?
- How can evolution occur at a microscopic and macroscopic level?
- Two couples want to know their risk of conceiving a child with cystic fibrosis. In one couple, neither partner has a family history of the disease; in the other, one partner knows he is a carrier. How do their risks differ?
- How does calculation of allele frequencies differ for an X-linked trait or disorder compared to one that is autosomal recessive?
- Why might understanding Hardy-Weinberg equilibrium be important in understanding epidemiology (the patterns of infectious diseases in populations)?
- Profiling of Y chromosome DNA implicated Thomas Jefferson in fathering a child of his slave, discussed in chapter 1. What might have been a problem with the conclusion?
- Glutaric aciduria type I (MIM 231680) causes progressive paralysis and brain damage. It is very common in the Amish of Lancaster County, Pennsylvania—0.25 percent of newborns have the disorder. What percentage of newborns are carriers for this condition?
- Torsion dystonia (MIM 128100) is a movement disorder that affects 1 in 1,000 Jewish people of eastern European descent (Ashkenazim). What is the carrier frequency in this population?
- The Finnish population has a 1 percent carrier frequency for a seizure disorder called myoclonus epilepsy (MIM 607876). Two people who have no relatives with the illness ask a genetic counselor to calculate the risk that they will conceive an affected child, based on their belonging to this population group. What is the risk?
- Maple syrup urine disease (MSUD) (see Reading 2.1) is autosomal recessive and causes mental and physical retardation, difficulty feeding, and a sweet odor to urine. In Costa Rica, 1 in 8,000 newborns inherits the condition. What is the carrier frequency of MSUD in this population?
- The amyloidoses are a group of inborn errors of metabolism in which sticky protein builds up in certain organs. Amyloidosis caused by a mutation in the gene encoding a blood protein called transthyretin (MIM 176300) affects the heart and/or nervous system. It is autosomal recessive. In a population of 177 healthy African Americans, four proved, by blood testing, to have one mutant allele of the transthyretin gene. What is the carrier frequency in this population?
- Ability to taste phenylthiocarbamide (PTC) (MIM 607751) is mostly determined by the gene *PTC*. The letters *T* and *t* are used here to simplify analysis. *TT* individuals taste a strong, bitter taste; *Tt* people experience a slightly bitter taste; *tt* individuals taste nothing.

A fifth-grade class of 20 students tastes PTC that has been applied to small pieces of paper, rating the experience as “very yucky” (*TT*), “I can taste it” (*Tt*), and “I can’t taste it” (*tt*). For homework, the students test their parents, with these results:

Of 6 *TT* students, 4 have 2 *TT* parents; and two have one parent who is *TT* and one parent who is *Tt*.

Of 4 students who are *Tt*, 2 have 2 parents who are *Tt*, and 2 have one parent who is *TT* and one parent who is *tt*.

Of the 10 students who can’t taste PTC, 4 have 2 parents who also are *tt*, but 4 students have one parent who is *Tt* and one who is *tt*. The remaining 2 students have 2 *Tt* parents.

Calculate the frequencies of the *T* and *t* alleles in the two generations. Is Hardy-Weinberg equilibrium maintained, or is this gene evolving?

Web Activities

14. Go to one of the biobank websites and describe a medical test or treatment that may be developed from its data.

Case Studies and Research Results

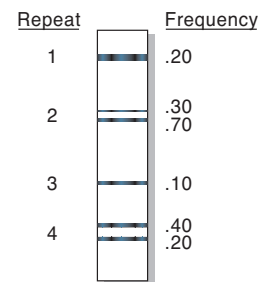
15. An extra row of eyelashes is an autosomal recessive trait that occurs in 900 of the 10,000 residents of an island in the south Pacific. Greta knows that she is a heterozygote for this gene, because her eyelashes are normal, but she has an affected parent. She wants to have children with a homozygous dominant man, so that the trait will not affect her offspring. What is the probability that a person with normal eyelashes in this population is a homozygote for this gene?

Forensics Focus

16. DNA dragnets have been so successful in catching criminals in several countries that some people in law enforcement have suggested storing DNA samples of everyone at birth, so that a DNA profile could be obtained from anyone at any time. Do you think that this is a good idea or not? Cite reasons for your answer.
17. On December 5, 1984, Theresa Fusco was raped and strangled near a roller-skating rink on Long Island, New York. Two similar crimes had occurred in previous months. Three young men were charged with the crime and then convicted, but proclaimed their innocence, maintaining that their confessions had been coerced and witnesses had lied. At their trial in 1990, defense lawyers requested DNA profiling, but the judge ruled that the technology was too unproven to use. In 2003, the case was reopened. Stored semen was taken from the “rape kit” and DNA profiled, leading to exoneration. The men had not killed Theresa Fusco but they had spent more than a decade in prison.
- Why might the judge have refused to consider DNA testing in 1990?
 - List the types of cells that could have been used to settle this case.
 - What information on the three suspects would be needed to interpret DNA patterns?
 - Do you think it is fair to decide whether or not a science-based forensic test or tool can be used based on how well a judge, jury, lawyers, or the public—who may have little or no training in genetics—understands how it works?
 - In 1992, lawyers Barry Scheck and Peter Neufeld, of the Cardozo School of Law in New York City, founded the nonprofit Innocence Project, described in the chapter opener. Consult the Innocence Project website, click on “Case Profiles,” select a case, and describe how DNA evidence exonerated a prisoner.
18. “Indirect genetic kinship analysis” is routinely used in forensic investigations of natural or manmade disasters. It uses a DNA profile for one person to lead to identification of a blood relative. For example, if 11 of the 13 STR alleles typically examined closely matched for a man whose brother was missing at the World Trade Center and a bone bit found there, the bone was assumed to have been from the brother. This approach is also being tried in criminal investigations. If a “cold hit” leads to a prisoner who couldn’t possibly have committed a crime (perhaps because he was locked up at the time), detectives investigate his closest relatives. Do you think that this is an ethical use of DNA profiling? What are the pros and cons of this approach?
19. Rufus the cat was discovered in a trash can by his owners, his body covered in cuts and bite marks and bits of gray fur clinging to his claws—gray fur that looked a lot like the coat of Killer, the huge, aptly-named hound next door. Fearful that Killer might attack their other felines, Rufus’ distraught owners brought his body to a vet, demanding a forensic analysis. The vet suggested that the hair might have come from a squirrel, but agreed to send appropriate samples to a veterinary genetic testing laboratory.

Identify the samples that the vet might have sent, and what information each could contribute to the case. (P.S.: This is a real case. Killer—not his real name—was found guilty on the basis of DNA testing, but was not punished because there were no human eyewitnesses. Rufus’ angry owners installed a fence.)

20. In a true crime that took place in Israel, a man knocked a woman unconscious with a cement block and then raped her. He was careful not to leave any hairs at the crime scene. But he left behind eyeglasses with unusual frames, and an optician helped police locate him. The man also left a half-eaten lollipop at the scene. DNA from blood taken from the suspect matched DNA from cheek-lining cells collected from the base of the telltale lollipop at four repeat loci on different chromosomes. Allele frequencies from the man’s ethnic group in Israel are listed beside the profile pattern below:



- For which of the tested repeats is the person a homozygote? How do you know this?
 - What is the probability that the suspect’s DNA matches that of the lollipop rapist by chance? (Do the calculation.)
 - The man’s population group is highly inbred—many people have children with relatives. How does this information affect the accuracy or reliability of the DNA profile? (P.S.—He was so frightened by the DNA analysis that he confessed!)
21. Simone, a recent college graduate, was spending 2 years using her biology degree to work in a clinic in Burma, in a town called Laputta. A devastating monsoon sent 18-foot waves over many of the neighboring villages, and survivors poured into the clinic. It had been raining for 8 days straight. Suddenly, a rush of water overwhelmed the clinic, taking equipment and children in its path. Simone tried to save the children but she, too, was washed away.
- Simone’s family in the United States was frantic, especially when her body was never found. Her father, a geneticist, insisted that investigators attempt to identify Simone’s DNA among evidence of human remains collected at the clinic. Explain how this might be done.
22. The governor of New York has proposed that DNA samples be taken from all people suspected of having committed a crime. What are the possible benefits and risks of such a program?



The ability to digest lactose (milk sugar) became more prevalent in populations after agriculture introduced dairy foods—thanks to evolution.

CHAPTER

15

Changing Allele Frequencies

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15.2 Migration

15.3 Genetic Drift

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Artificial Selection

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Evolving HIV

Balanced Polymorphism

15.6 Putting It All Together: PKU

15.7 Eugenics

The Evolution of Lactose Tolerance

For millions of people who have lactose intolerance, dairy food causes cramps, bloating, gas, and diarrhea. They no longer produce lactase, an enzyme made in early childhood that breaks down the milk sugar lactose into easily digested sugars. People who can digest dairy have lactase persistence (MIM 223000). Four genes control the ability to digest milk sugar. Different populations have different proportions of lactose-intolerant versus lactase-persistent people. Clues in DNA suggest why: agriculture.

As dairy farming spread around the world, people who had gene variants enabling them to digest milk into adulthood had an advantage, and had more children. Over time, populations that consumed dairy foods had more people with lactase persistence. In contrast, in populations with few or no dairy foods, lactose intolerance was not a problem, and so those gene variants persisted.

The link between lactose intolerance and agriculture is why today, the European American population only has 10 percent lactose intolerance, whereas among Asian Americans, who eat far less dairy, 90 percent have lactose intolerance. Seventy-five percent of African Americans and Native Americans have lactose intolerance.

15.1 Nonrandom Mating

Historically, we seem to have gone out of our way to ensure that the very specific conditions necessary for Hardy-Weinberg equilibrium—unchanging allele frequencies from generation to generation—do not occur, at least for some genes. Religious restrictions and personal preferences guide our choices of mates. Wars and persecution kill certain populations. Economic and political systems enable some groups to have more children. We travel, shuttling genes in and out of populations. Natural disasters and new diseases reduce populations to a few individuals, who then rebuild their numbers, at the expense of genetic diversity. These factors, plus mutation and a reshuffling of genes at each generation, make a gene pool very fluid.

The ever-present and interacting forces of nonrandom or selective mating, migration, genetic drift, mutation, and natural selection work to differing degrees to shape populations at the allele level. Changing allele frequencies can change genotype frequencies—which in turn can change phenotype frequencies. In a series of illustrations throughout this chapter, colored shapes represent individuals who have specific genotypes. Figure 15.14 then combines the illustrations to summarize the

chapter. We begin our look at the forces that change allele frequencies in populations with nonrandom mating.

In the theoretical state of Hardy-Weinberg equilibrium, individuals of all genotypes are presumed equally likely to mate and to choose partners at random. For some traits this is true—we do not choose partners based on blood type, for example—but other traits do influence our mate choices. We choose partners based on physical appearance, ethnic background, intelligence, and shared interests. We marry people similar to ourselves about 80 percent of the time (**figure 15.1**). Worldwide, about one-third of all marriages occur between people who were born fewer than ten miles apart! This nonrandom mating is a major factor in changing allele frequencies in human populations.

Nonrandom mating occurs when certain individuals contribute more to the next generation than others (**figure 15.2**). This is common in agriculture when semen from one prize bull is used to inseminate thousands of cows, or a field of genetically identical crops is planted. Such an extreme situation can arise in a human population when a man fathers many children. A striking mutation can reveal such behavior. In the Cape population of South Africa, for example, a Chinese immigrant known as Arnold had a very rare dominant mutation that



Figure 15.1 Partner choice affects genetic diversity in populations. Most of us marry people from the same population group who are, therefore, genetically similar to ourselves.

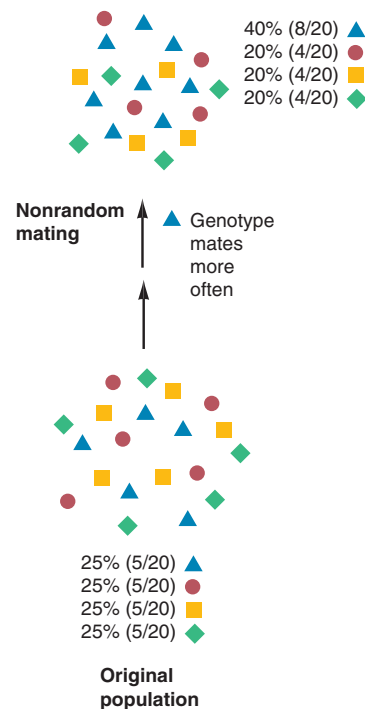


Figure 15.2 Nonrandom mating alters allele frequencies. The different-colored shapes represent individuals with distinctive genotypes. If Hardy-Weinberg equilibrium exists for these genes in this population, then the percentages will remain the same through the generations. However, the blue triangle genotype is more reproductively successful, skewing the allele frequencies in the next generation.

causes teeth to fall out before age 20. Arnold had seven wives. Of his 356 living descendants, 70 have the dental disorder. The frequency of this allele in the Cape population is exceptionally high, thanks to Arnold.

The high frequency of autosomal recessive albinism among Arizona's Hopi Indians also reflects nonrandom mating. Albinism is uncommon in the general U.S. population, but it affects 1 in 200 Hopi Indians. The reason for the trait's prevalence is cultural—men with albinism often stay back and help the women, rather than risk severe sunburn in the fields with the other men. They contribute more children to the population because they have more contact with the women.

The events of history reflect nonrandom mating patterns. When a group of people is subservient to another, genes tend to “flow” from one group to the other as the males of the ruling class have children with females of the underclass—often forcibly. Historical records and DNA sequences show this directional gene flow phenomenon. For example, Y chromosome analysis suggests that Genghis Khan, a Mongolian warrior who lived from 1162 to 1227, had sex with so many women that today, 1 in every 200 males living between Afghanistan and northeast China shares his Y—that's 16 million men (**figure 15.3**)!

Despite our partner preferences, many traits do mix randomly in the next generation. This may be because we are unaware of these characteristics or because we do not consider them in choosing partners. In populations where AIDS is



Figure 15.3 A prevalent Y. Genghis Khan left his mark on many male descendants in the form of his Y chromosome. Rape of women on a sweeping scale spread the chromosome in certain Asian populations.

extremely rare or nonexistent, for example, the two mutations that render a person resistant to HIV infection are in Hardy-Weinberg equilibrium. This would change, over time, if HIV arrives, because the people with these mutations would become more likely to survive to produce offspring—some of whom would perpetuate the protective mutation. Natural selection would intervene, ultimately altering allele frequencies.

Many blood types are in Hardy-Weinberg equilibrium because we do not choose partners by blood type. Yet sometimes the opposite occurs. People with mutations in the same gene meet when their families participate in programs for people with the associated disorder. For example, more than two-thirds of relatives visiting a camp for children with cystic fibrosis are likely to be carriers, compared to the 1 in 23 or fewer in large population groups.

People can avoid genetic disease with controlled reproduction. In a program in New York City called Dor Yeshorim, for example, young people take tests for a dozen genetic disorders that are much more common among Jewish people of eastern European descent (Ashkenazim). Results are stored in a confidential database. Two people wishing to have children together can find out if they are carriers for the same disorder. If so, they may elect not to have children. Thousands of people have been tested, and the program is partly responsible for the near-disappearance of Tay-Sachs disease among Ashkenazi Jews. The very few cases each year are usually in non-Jews, because they have not been tested.

A population that practices consanguinity has very nonrandom mating. Recall from chapter 4 that in a consanguineous relationship, “blood” relatives have children together. On the family level, this practice increases the likelihood that harmful recessive alleles from shared ancestors will be combined and passed to offspring, causing disease. The birth defect rate in offspring is 2.5 times the normal rate of about 3 percent. On a population level, consanguinity decreases genetic diversity. The proportion of homozygotes rises as that of heterozygotes falls.

Some populations encourage marriage between cousins, which increases the incidence of certain recessive disorders. In certain parts of the middle east, Africa, and India, 20 to 50 percent of marriages are between cousins, or uncles and nieces. The tools of molecular genetics can reveal these relationships. Researchers traced DNA sequences on the Y chromosome and in mitochondria among residents of an ancient, geographically isolated “micropopulation” on the island of Sardinia, near Italy. They consulted archival records dating from the village's founding by 200 settlers around 1000 A.D. to determine familial relationships. Between 1640 and 1870, the population doubled, reaching 1,200 by 1990. Fifty percent of the present population descends from just two paternal and four maternal lines, and 86 percent of the people have the same X chromosome. Researchers are analyzing disorders that are especially prevalent in this population, which include hypertension and a kidney disorder.

Worldwide, about 960 million married couples are related, and know of their relationship. Also contributing to nonrandom mating is endogamy, which is marriage within a community. In an endogamous society, spouses may be distantly related and unaware of the connection.

Key Concepts

1. People choose mates for many reasons, and they do not contribute the same numbers of children to the next generation. These practices change allele frequencies in populations.
2. Traits lacking obvious phenotypes may be in Hardy-Weinberg equilibrium.
3. Consanguinity and endogamy increase the proportion of homozygotes in a population.

15.2 Migration

Large cities, with their pockets of ethnicity, defy Hardy-Weinberg equilibrium by their very existence. Waves of immigrants formed the population of New York City, for example. The original Dutch settlers of the 1600s had different alleles than those in today's metropolis of English, Irish, Slavics, Africans, Hispanics, Italians, Asians, and many others. **Figure 15.4** depicts the effect on allele and genotype frequencies when individuals join a migrating population. Clues to past migrations lie in historical documents as well as in differing allele frequencies in regions defined by geographical or language barriers.

The frequency of the allele that causes galactokinase deficiency (MIM 230200) in several European populations reveals how people with this autosomal recessive disorder migrated (**figure 15.5**). Galactokinase deficiency causes cataracts (clouding of the lens) in infants. It is very common among a population of 800,000 gypsies, called the Vlach Roma, who live in Bulgaria. It affects 1 in 1,600 to 2,500 people among them, and 5 percent of the people are carriers. But among all

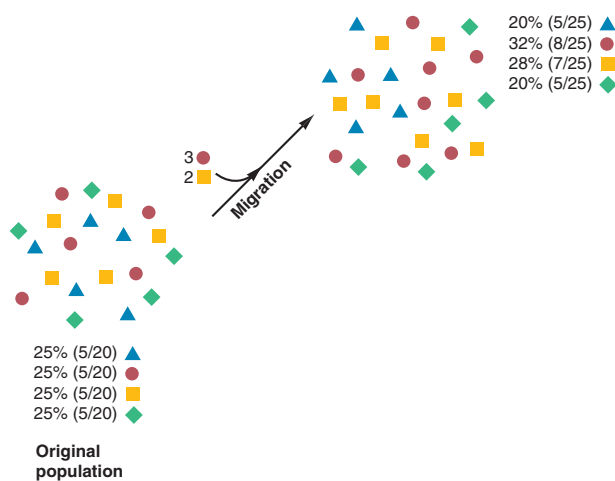


Figure 15.4 Migration alters allele frequencies. If the population travels and picks up new individuals, allele (and genotype) frequencies can change.

gypsies in Bulgaria as a whole, the incidence drops to 1 in 52,000. As the map in figure 15.5 shows, the disease becomes rarer to the west. This pattern may have arisen when people with the allele settled in Bulgaria, with only a few individuals or families moving westward.

Allele frequencies often reflect who ruled whom. For example, the frequency of ABO blood types in certain parts of the world today mirrors past Arab rule. The distribution of ABO blood types is very similar in northern Africa, the Near East, and southern Spain. These are precisely the regions where Arabs ruled until 1492. The uneven distribution of allele frequencies can also reveal when and where nomadic peoples stopped. For example, in the eighteenth century, European Caucasians called trekboers migrated to the Cape area of South Africa. The men stayed and had children with the native women of the Nama tribe. The mixed society remained fairly isolated, leading to the distinctive allele frequencies found in the present-day people of color of the area.

Sometimes allele frequencies change from one neighboring population to another. This phenomenon is termed a **cline**. Changing allele frequencies usually reflect migration patterns, as immigrants introduced alleles and emigrants removed them. Clines may be gradual, reflecting unencumbered migration paths, but barriers often cause more abrupt changes in allele frequencies. Geographical formations such as mountains and bodies of water may block migration, maintaining population differences in allele frequencies on either side of the barrier. Language differences may also isolate alleles, if people who cannot communicate tend not to have children together.

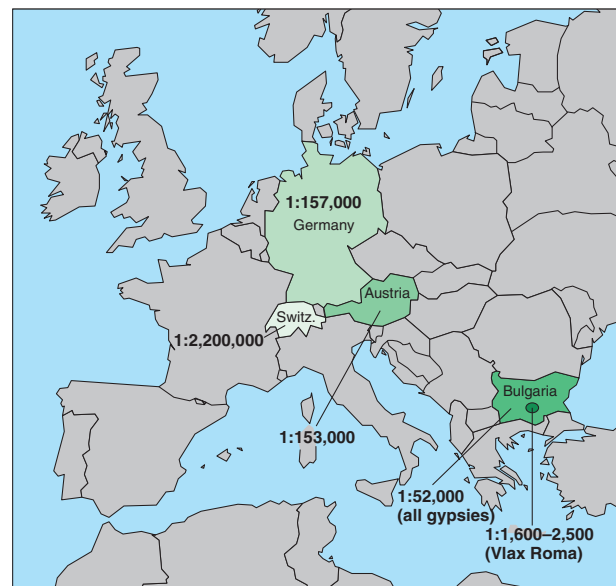


Figure 15.5 Galactokinase deficiency in Europe illustrates a cline. This autosomal recessive disorder that causes blindness varies in prevalence across Europe. It is most common among the Vlach Roma gypsies in Bulgaria. The condition becomes much rarer to the west, as indicated by the shading from dark to light green.

Allele frequencies up and down the lush strip of fertile land that hugs the Nile River illustrate the concept of clines. Researchers found a gradual change in mitochondrial DNA sequences in 224 people who live on either side of the Nile, an area settled 15,000 years ago. The farther apart two individuals live along the Nile, the less alike their mtDNA. This is consistent with evidence from mummies and historical records that indicate the area was once kingdoms separated by wars and language differences. If the area had been one large interacting settlement, then the DNA sequences would have been more mixed. Instead, the Nile may have served as a “genetic corridor” between Egypt and sub-Saharan Africa.

Comparing clines is also consistent with geography and history. For example, agriculture spread faster from east to west in Eurasia because latitude does not change much. As a result, there is less evidence of gene flow in this relatively unchanging environment—even if people migrated, they tended to have the same gene variants as the people had in their new homes. In the Americas, the spread of agriculture was north to south. Differing hours of daylight and climate with longitude presented different environmental challenges, which are reflected in greater genetic change from north to south than from east to west.

Clines may also reflect human dependence on communication. Geneticists correlated twenty blood types to geographically defined regions of Italy and to areas where a single dialect is spoken. They chose Italy because it is rich in family history records and linguistic variants. Six of the blood types varied more consistently with linguistically defined subregions than with geographical regions. Perhaps differences in language prevent people from socializing, keeping alleles within groups.

Key Concepts

1. Migration alters genotype frequencies by adding and removing alleles from populations.
2. Clines are gradual changes in allele frequencies between neighboring populations.
3. Geographical barriers and language differences often create great differences in allele frequencies.

15.3 Genetic Drift

When a small group separates from a larger population, or reproduces only among themselves, allele frequencies may change as a result of chance sampling from the whole (**figure 15.6**). This change in allele frequency that occurs when a small group separates from the larger whole is termed **genetic drift**. It is like reaching into a bag of jellybeans and, by chance, grabbing only green and yellow ones. The allele frequency changes in genetic drift are random and unpredictable, just as reaching into the jellybean bag a second time might yield mostly black and orange candies.

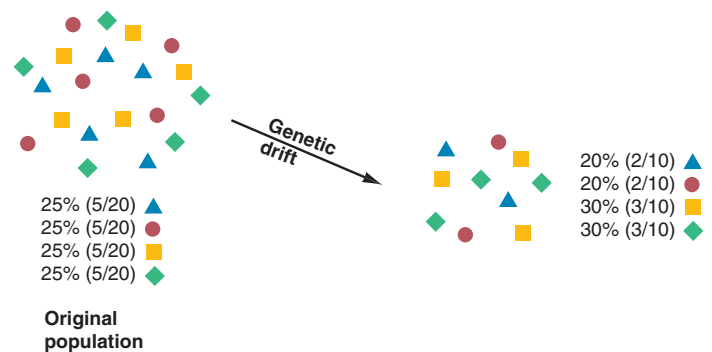


Figure 15.6 Genetic drift alters allele frequencies. If members of a population leave or do not reproduce, allele frequencies can change by chance sampling of a small population. When half of this population does not contribute to the next generation, two genotypes increase in frequency and two decrease.

Genetic drift occurs when the population size plummets, due either to migration, to a natural disaster or geographic barrier that isolates small pockets of people, or to the consequences of human behavior. Members of a small community might reproduce only among themselves, which keeps genetic variants within their ethnic group. For example, the skin-lightening condition vitiligo (MIM 193200) is much more common in a small community isolated in the mountains of northern Romania than elsewhere in the nation. Genetic drift is also seen within large cities. Pittsburgh, Pennsylvania and New York City are more mosaics of groups with distinct ethnic flavors than “melting pots” of mixed heritage.

Some groups of people become isolated in several ways—geographically, linguistically, and by choice of partners. Such populations often have a high incidence of several otherwise rare inherited conditions. The native residents of the Basque country in the western part of the Pyrenees Mountains between France and Spain, for example, still speak remnants of Euskera, a language the first European settlers brought in more than 10,000 years ago. The Basques have unusual frequencies of certain ABO and Rh blood types, rare mtDNA sequences and cell surface antigen patterns, and a high incidence of a mild form of muscular dystrophy.

The Founder Effect

A common type of genetic drift in human populations is the **founder effect**, which occurs when small groups leave home to found new settlements. The new colony may have different allele frequencies than the original population.

Founder populations can amplify certain alleles. This shows up in increased disease frequencies. Consider an isolated community of about 10,000 members of the Church of Jesus Christ of Latter Day Saints, who live on a plateau on the Utah-Arizona border. Half of all known cases of fumarate deficiency (MIM 606812), which causes mental retardation, seizures, and coma, occur in this area. Nearly 80 percent of the population descends from two original settlers. Widespread

Population	Number of Founders	Number of Generations	Population Size Today
Costa Rica	4,000	12	2,500,000
Finland	500	80–100	5,000,000
Hutterites	80	14	36,000
Japan	1,000	80–100	120,000,000
Iceland	25,000	40	300,000
Newfoundland	25,000	16	500,000
Quebec	2,500	12–16	6,000,000
Sardinia	500	400	1,660,000

cousin-cousin marriages as well as polygamy (men with multiple wives) and social isolation have joined the powerful founder effect to keep the disease in the population. In 2008, the U.S. government removed many children from this community on suspicion of abuse. Some of the children returned a year later—but some chose to remain in the outside world. **Table 15.1** lists some other founder populations.

A powerful founder effect appears in the French Canadian population of Quebec. Their lack of diversity in disease-causing mutations reflects a long history of isolation. Consider breast cancer caused by the *BRCA1* gene. More than 500 alleles are known worldwide, yet only four are seen among French Canadians. Several inborn errors of metabolism are also more common in this group. The French Canadians have ideal characteristics for gene discovery: many generations since founding (14), few founders (about 2,500), a high rate of population expansion (74 percent increase per generation), a large present-day population (about 6 million), and most marriages within the group.

The French Canadian population exemplifies genetic drift because the people have kept mostly to themselves within a larger population. The French founded Quebec City in 1608. Until 1660, the population grew as immigrants arrived from France, and then began to increase from births. More than 10,000 French had arrived by the time the British took over in 1759, but many of them had headed westward. In Quebec, religious, language, and other cultural differences separated the French and English gene pools. The French Canadian population of Quebec grew from the 2,500 or so founding genotypes to about 6 million individuals today.

The cultural and physical isolation in Canada created an unusual situation—a founder effect within a founder effect. In the nineteenth century, when agricultural lands opened up about 150 miles north of Quebec, some families migrated north. Their descendants, who remained in the remote area, form an incredibly genetically homogeneous subpopulation of founders split off from the original set of founders.

A classic example of a founder effect within a larger population is the Dunker community of Germantown, Pennsylvania. Excellent historical records combined with distinctive traits enabled geneticists to track genetic drift from the larger surrounding population. The Dunkers came from Germany between 1719 and 1729, but they have lived among others since that time. Still, the frequencies of some genotypes are different among the Dunkers than among their non-Dunker neighbors, and they are also different from the frequencies seen among people living in their native German village. The Dunkers have a different distribution of blood types (**table 15.2**) and much higher incidence of attached earlobes, hyperextensible thumbs, hairs in the middle of their fingers, and left-handedness compared to the other two groups.

Founder effects can be studied at the phenotypic and genotypic levels. Phenotypically, a founder effect is indicated when a community of people, known from local history to have descended from a few founders, have inherited traits and illnesses that are rare elsewhere. This is striking among the Old Order Amish and Mennonites of Lancaster County, Pennsylvania. Often, worried parents would bring their ill children to medical facilities in Philadelphia. Over the years, researchers realized that these people are subject to an array of extremely rare conditions (**table 15.3** and **figure 15.7**). For example, Victor McKusick, founder of *Online Mendelian Inheritance in Man*, discovered and described cartilage-hair hypoplasia after six Amish children died at a Philadelphia hospital from chickenpox in 1965. Part of their inherited syndrome was impaired immunity. Until McKusick made the connection, the other symptoms—including dwarfism, sparse hair, and anemia—were not recognized as part of a syndrome. Today, as many geneticists study inherited diseases common among the Amish and Mennonites, treatments are becoming available, from special diets to counter inborn errors of metabolism to gene therapy.

Blood Type	Population		
	U.S.	Dunker	European
ABO System			
A	40%	60.0%	45%
B, AB	15%	5.0%	15%
Rh ⁻	15%	11.0%	15%
MN System			
M	30%	44.5%	30%
MN	50%	42.0%	50%
N	20%	13.5%	20%

Table 15.3 Inherited Conditions Common Among the Amish and Mennonites of Lancaster County, Pennsylvania

Disorder	MIM	Signs and Symptoms (Phenotype)
Ataxia telangiectasia	208900	Increased sensitivity to radiation, loss of balance and coordination, red marks on face, delayed sexual maturation, lung infections, diabetes, high risk of cancer
Bipolar affective disorder	Several	Mood swings (manic depression)
Cartilage-hair hypoplasia (metaphyseal chondrodysplasia, McKusick type)	250250	Dwarfism, sparse hair, anemia, poor immunity
Crigler-Najjar syndrome	218800	Bilirubin buildup, jaundice, brain damage
Ellis-van Creveld syndrome	225500	Dwarfism, short fingers, underdeveloped nails, polydactyly, hair "blaze" pattern, heart disease, fused bones, teeth at birth
Glutaric aciduria type I	231670	Paralysis, brain damage
Homocystinuria	236200	Damaged blood vessels, stroke, heart attack
Limb-girdle muscular dystrophy	253600	Progressive muscle weakness in limbs
Maple syrup urine disease	248600	Sweet-smelling urine, sleepiness, vomiting, mental retardation
Metachromatic leukodystrophy	250100	Rigid muscles, convulsions, mental deterioration
Morquio syndrome	252300	Clouded corneas, abnormal skeleton and aortic valve
Sudden infant death syndrome with dysgenesis of testes	608800	Sudden cessation of heartbeat and breathing; underdeveloped testes



Figure 15.7 Ellis-van Creveld syndrome. This Amish child has autosomal recessive Ellis-van Creveld syndrome, which causes dwarfism, extra fingers, heart disease, fused wrist bones, and teeth at birth. Seven percent of the people of this community have the mutant allele. Heterozygotes have the milder condition Weyers acrocentric dysostosis.

In addition to historical records, differences in allele frequencies in a smaller population compared to those in the general population suggest a founder effect. The incidence of certain diseases in Lancaster County, for example, is astounding. Maple syrup urine disease affects 1 in 225,000 newborns in the United States, but 1 in 400 newborns among the Lancaster families! A research fellow at Children's Hospital in Philadelphia discovered that cerebral palsy in several young children from Lancaster County attributed to oxygen deprivation at birth was actually an inborn error of metabolism called glutaric aciduria type I. He went from farm to farm, tracking cases against genealogical records, and found that *every family* that could trace its roots back to the founders had members who had the disease! Today, 1 in 200 newborns in this population have the condition.

A mutation that is the same in all affected individuals in a population is strong evidence of a founder effect due to descent from shared ancestors. The Bulgarian gypsies who have galactokinase deficiency, for example, all have a mutation that is extremely rare elsewhere. In contrast, a population with several mutations that cause the same disorder is more likely to have picked up those variants from people joining the group, rather than from descent from shared founders.

Very often when a disease-associated allele is identical in DNA sequence among people in the same population, so is the DNA surrounding the gene. This pattern indicates that a portion of a chromosome, rather than just the disease-causing gene, has been passed among the members of the population from its

founders. For this reason, many studies that trace founder effects examine haplotypes that include tightly linked genes.

When historical or genealogical records are particularly well kept or recent, founder effects can sometimes be traced to the very beginning. This is the case for the Afrikaner population of South Africa. The 2.5 million Afrikaners descended from a small group of Dutch, French, and German immigrants who had huge families, often with as many as ten children. In the nineteenth century, some Afrikaners migrated northeast to the Transvaal Province, where they lived in isolation until the Boer War in 1902 introduced better transportation.

Today, 30,000 Afrikaners have porphyria variegata (see figures 5.5 and 5.6). All affected people descended from one couple who came from the Netherlands in 1688! Today's allele frequency in South Africa is far higher than that in the Netherlands because the founding couple had many children—who, in turn, had large families, passing on and amplifying the dominant mutation.

Founder effects are also evident in more common illnesses, where populations have different mutations in the same gene. *BRCA1* breast cancer, for example, is most prevalent among Ashkenazi Jewish people. Nearly all affected individuals have the same 3-base deletion. In contrast, *BRCA1* breast cancer is rare in blacks, but it affects families from the Ivory Coast in Africa, the Bahamas, and the southeastern United States. They share a 10-base deletion, probably inherited from West Africans ancestral to all three modern groups. Slaves brought the disease to the United States and the Bahamas between 1619 and 1808, but some of their relatives who stayed in Africa have perpetuated the mutant allele there.

Population Bottlenecks

A **population bottleneck** occurs when many members of a group die, and only a few are left to replenish the numbers. The new population has only those alleles in the small group that survived the catastrophe. An allele in the replenished population might become more common in the replenished population than it was in the original larger group. Therefore, the new population has a much more restricted gene pool than the larger ancestral population, with some variants amplified, others diminished.

Population bottlenecks can occur when people (or other animals) colonize islands. An extreme example is seen among the Pingelapese people of the eastern Caroline Islands in Micronesia. Four to 10 percent are born with “Pingelapese blindness,” an autosomal recessive combination of colorblindness, nearsightedness, and cataracts also called achromatopsia (MIM 603096). Elsewhere, only 1 in 20,000 to 50,000 people inherits the condition. Nearly 30 percent of the Pingelapese are carriers. The prevalence of the blindness among the Pingelapese stems from a typhoon in 1780 that killed all but nine males and ten females who founded the present population. This severe population bottleneck, plus geographic and cultural isolation, increased the frequency of the blindness gene as the population resurged.

A more widespread population bottleneck occurred as a consequence of the early human expansion from Africa, discussed in chapter 16. As numbers dwindled during the journeys and then were replenished as people settled down, mating among relatives led, over time, to an increase in homozygous recessive genotypes compared to ancestral populations that maintained their genetic diversity in Africa. These bottlenecks are reflected today in the persistence of genetic diversity among African populations. The lack of genetic diversity in some modern human populations is evident as “runs of homozygosity,” which are chromosome regions that vary little from person to person. Runs of homozygosity generally represent regions that are inherited from shared ancestors. They are common, for example, in highly purebred dogs.

Figure 15.8 illustrates schematically the dwindling genetic diversity that results from a population bottleneck. Today's cheetahs live in just two isolated populations of a few thousand animals in South and East Africa. Their numbers once exceeded 10,000. The South African cheetahs are so alike genetically that even unrelated animals can accept skin grafts from each other. Researchers attribute the cheetahs' genetic uniformity to two bottlenecks—one that occurred at the end of the most recent ice age, when habitats changed, and another following mass slaughter by humans in the nineteenth century. However, the good health of the animals today indicates that the genes that have survived enable the cheetahs to thrive in their environment.

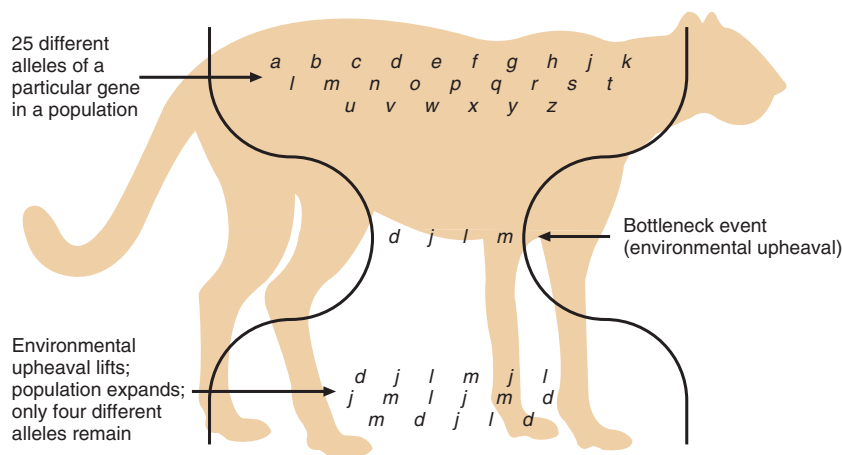


Figure 15.8 Population bottlenecks. A population bottleneck occurs when the size of a genetically diverse population falls, remains at this level for a time, and then expands again. The new population loses some genetic diversity if alleles are lost.

Table 15.4 Autosomal Recessive Genetic Diseases Prevalent Among Ashkenazi Jewish Populations

Disorder	MIM	Signs and Symptoms (Phenotype)	Carrier Frequency
Bloom syndrome	210900	Sun sensitivity, short stature, poor immunity, impaired fertility, increased cancer risk	1/110
Breast cancer	113705, 600185	Malignant breast tumor caused by mutant <i>BRCA1</i> or <i>BRCA2</i> genes	3/100
Canavan disease	271900	Brain degeneration, seizures, developmental delay, early death	1/40
Familial dysautonomia	223900	No tears, cold hands and feet, skin blotching, drooling, difficulty swallowing, excess sweating	1/32
Gaucher disease	231000	Enlarged liver and spleen, bone degeneration, nervous system impairment	1/12
Niemann-Pick disease type A	257200	Lipid accumulation in cells, particularly in the brain; mental and physical retardation, death by age three	1/90
Tay-Sachs disease	272800	Brain degeneration causing developmental retardation, paralysis, blindness, death by age four	1/26
Fanconi anemia type C	227650	Deficiencies of all blood cell types, poor growth, increased cancer risk	1/89

Human-wrought disasters that kill many people can cause population bottlenecks—perhaps even more severely, because aggression is typically directed at particular groups, while a typhoon indiscriminately kills anyone in its path. The Chmielnicki massacre was one of many attacks against the Ashkenazim. Overall, these acts have left a legacy of several inherited diseases that are at least ten times more common among Jewish people than in other populations (**table 15.4**), although some of the disorders have become rarer as genetic testing has become available.

The Chmielnicki massacre began in 1648, when a Ukrainian named Bogdan Chmielnicki led a massacre against the Polish people, including peasants, nobility, and the Jewish people, in retaliation for a Polish nobleman's seizure of his possessions. By 1654, Russians, Tartars, Swedes, and others joined the Ukrainians in wave after wave of violence against the Polish people. Thousands perished, with only a few thousand Jewish people remaining.

The Jewish people have survived many massacres, and therefore many population bottlenecks; after the Chmielnicki massacre, like the others, their numbers grew again. From 1800 to 1939, the Jewish population in Eastern Europe swelled to several million. Yet massacres continued. Jewish people also tended to have children only with each other. Both of these factors—nonrandom mating and population bottlenecks—changed allele frequencies and contributed to the high incidence of certain inherited diseases seen among the Ashkenazim today. Several genetic testing companies offer “Jewish genetic disease” panels that are not meant to discriminate or stereotype, but are based on a genetic fact of life—some illnesses are more common in certain populations.

Key Concepts

1. Genetic drift occurs when a subset of a population has different allele frequencies than the larger population.
2. The founder effect occurs when a few individuals leave a community to start a new settlement. The resulting population may, by chance, either lack some alleles from the original population or have high frequencies of others.
3. In a population bottleneck, many members die, and only a few contribute to the next generation.

15.4 Mutation

A major and continual source of genetic variation in populations is mutation—when one allele changes into another and the change is passed to offspring (**figure 15.9**). Genetic variability also arises from crossing over and independent assortment during meiosis, but these events recombine existing traits rather than introduce new ones.

If a DNA base change occurs in a part of a gene that encodes part of a protein necessary for its function, then an altered trait may result. Another way that genetic change can occur from generation to generation is in the numbers of repeats of copy number variants (CNVs). These function as alleles.

Natural selection, discussed in the next section, eliminates alleles that adversely affect reproduction. Yet harmful recessive alleles are maintained in heterozygotes and are reintroduced by new mutation. Therefore, all populations have some alleles that would be harmful if homozygous. The

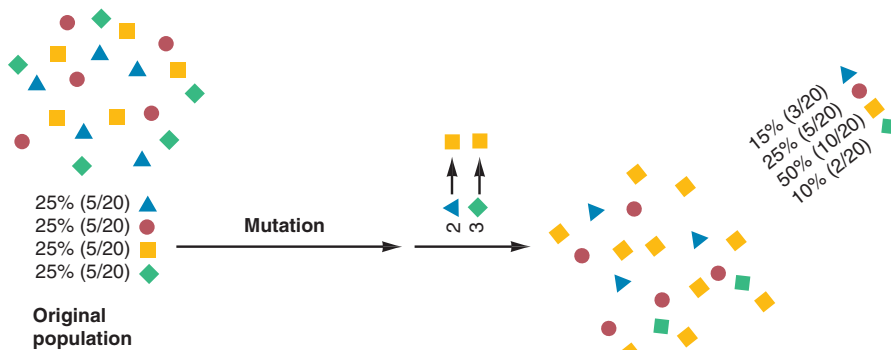


Figure 15.9 Mutation alters allele frequencies. If one allele changes into another from one generation to the next, genotype frequencies can change.

In natural selection, reproductive success is all-important, for this is what transmits favorable alleles and weeds out the unfavorable ones, ultimately impacting population structure and therefore microevolution. In the common phrase used synonymously with natural selection—“survival of the fittest”—“fit” actually refers to reproductive success, not to physical prowess or intelligence (unless those traits lead directly to reproductive success). In a Darwinian sense, an unattractive and out-of-shape parent of ten is more “fit” than a gorgeous triathlete with one child.

collection of such deleterious alleles in a population is called its **genetic load**.

The contribution that mutation makes to counter Hardy-Weinberg equilibrium is quite small compared to the influence of migration and nonrandom mating, because mutation is rare. Natural selection has the greatest influence. The spontaneous mutation rate is only about 30 bases per haploid genome in each gamete. Each of us probably has at least five “lethal equivalents”—alleles or allele combinations that if homozygous would kill us or make us too sick to have children.

Key Concepts

1. Mutation alters genotype frequencies by introducing new alleles.
2. Heterozygotes and new mutations maintain the frequencies of deleterious alleles in populations.

15.5 Natural Selection

Environmental change can alter allele frequencies when individuals with certain phenotypes are more likely to survive and reproduce than others. This differential survival to reproduce guided by environmental change is **natural selection** (figure 15.10). The chapter opener chronicles natural selection acting on the gene variant that enables people to digest lactose.

Inability to digest lactose is actually the wild type condition—it predominated before people began domesticating mammals and drinking their milk. Most of the mutations that introduced lactase persistence are point mutations. Another dietary illustration of natural selection involves CNVs. Populations that follow high-starch diets tend to have more copies of the gene that encodes salivary amylase, the digestive enzyme that begins to break down starch in the mouth. Members of populations that follow low-starch diets have fewer copies of the gene—and presumably less of the enzyme.

Negative and Positive Selection

Natural selection can retain an advantageous trait or banish one that has become dangerous in the prevailing environment. Retaining a trait is termed positive natural selection, and getting rid of a trait is termed negative natural selection, but the “positive” and “negative” are not value judgments—they merely refer to staying or leaving. In Darwin’s time natural selection was thought to be primarily negative, but the ability to sequence genes has enabled us to actually measure and track the instances of positive selection that have sculpted our differences from our closest primate relatives.

Subtle nuances in DNA sequence provide a signature of sorts for positive selection. Specifically, a sign of positive selection is a gene in humans that has a counterpart in other primates, but in humans has at least one distinctive difference in the amino acid sequence. A change in the DNA sequence that

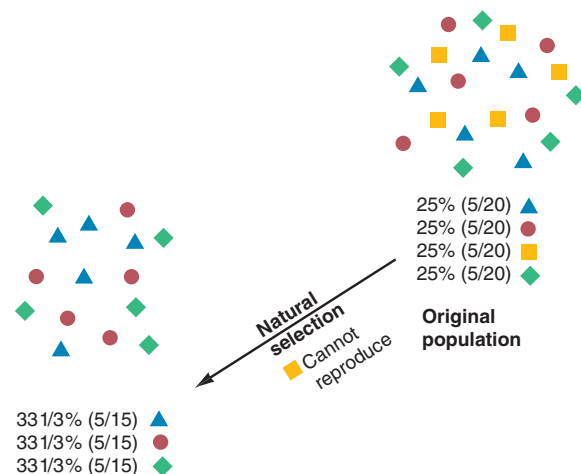


Figure 15.10 Natural selection alters allele frequencies. If health conditions impair the ability of individuals of a certain genotype to reproduce, allele frequencies can change.

does *not* substitute an amino acid does not change the protein, and therefore has no effect on the phenotype. Such a change therefore cannot be subject to natural selection.

Another sign of positive selection is a gene that varies little, if at all, from person to person. Positive selection is, in a sense, an evolutionary version of “if it ain’t broke, don’t fix it.” A study of 1.2 million SNPs across the genomes of African Americans, European Americans, and Chinese revealed 101 regions that have signs of positive selection. Genes in these regions take part in the sense of smell, digestion, nervous system development, immunity, muscle contraction, and response to stress. Reading 16.1 offers several examples of traits that were positively selected in human evolution.

Artificial Selection

Natural selection acts on preexisting genetic variants and is uncontrolled and largely unpredictable. In contrast, artificial selection is controlled breeding with the intent of perpetuating individuals with a particular phenotype, such as a crop plant or fancy pet. Darwin’s idea of natural selection actually grew from his observations of artificial selection of pigeons.

Our pets are products of intense artificial selection. Dalmatians, for example, had a mutation that caused painful bladder stones. The trait was bred out of the dogs by setting up specific crosses that preserved the coveted spots. **Figure 15.11** shows extremes of dog breeding. The ancestral dog was probably similar to the modern wild dog of Australia, the dingo. Dogs were introduced from southeast Asia in a tiny founder population about 40,000 years ago. People bred



Figure 15.11 Dogs small and large. It is hard to believe that the diminutive Chihuahua and the Great Dane are members of the same species! One gene that contributes to canine body size encodes insulin-like growth factor 1 (*IGF-1*). Two alleles control abundance of the encoded protein. Small breeds tend to have genotypes that limit the amount of the growth factor, whereas larger breeds as well as ancestral wolves have genotypes associated with abundant growth factor. However, the fact that the “low-IGF-1” genotype is seen in at least two large breeds—Mastiffs and Rottweilers—suggests that body weight in dogs is controlled by more than one gene, as is the case in humans (see section 7.3).

the diverse dog breeds of today which, thanks to extreme artificial selection, suffer from more than 300 inherited disorders. Would natural selection have led to the Pekinese with its unstable eyeballs, or the bulldog with its dental woes and notorious dog breath?

Interestingly, black wolves and coyotes picked up the mutation that darkens their fur by breeding with domesticated dogs, about 14,000 years ago when canines accompanied people across the Bering Strait land bridge to North America (see figure 16.21). The mutation is in a gene that encodes an immune system protein. Dark coat color may have increased in the wolf and coyote populations due to positive selection for better immunity. The trait is the first known example of artificial selection leading to positive selection.

Cats were first domesticated in the Near East when agriculture began, about 10,000 years ago. They descended from one of five subspecies of wildcats.

Tuberculosis Ups and Downs—and Ups

Natural selection can be seen in the appearance or reemergence of infectious diseases. If infection kills before reproductive age or impairs fertility, its spread will ultimately remove from the population individuals susceptible to infection. Disease incidence falls as only survivors are left. But if conditions change, the disease may resurge. This has happened with tuberculosis (TB).

When TB first appeared in the Plains Indians of the Qu’Appelle Valley Reservation in Saskatchewan, Canada in the mid-1880s, it struck swiftly and lethally, infecting many organs. Ten percent of the population died. But by 1921, TB tended to affect only the lungs, and only 7 percent of the population died annually from it. By 1950, mortality was down to 0.2 percent. Some people were symptomless carriers.

Outbreaks of TB ran similar courses in other human populations. The disease appeared in crowded settlements where the bacteria easily spread in exhaled droplets. In the 1700s, TB raged through the cities of Europe. Immigrants brought it to the United States in the early 1800s, where it also swept the cities. But TB incidence and virulence fell dramatically in the cities of the industrialized world in the first half of the twentieth century—before antibiotic drugs were discovered. What tamed tuberculosis?

Natural selection, operating on both the bacterial and human populations, lessened the virulence of the infection. Some people inherited resistance and passed this beneficial trait on. At the same time, the most virulent bacteria killed their hosts so quickly that the victims had no time to spread the infection. As the deadliest bacteria were selected out of the population (negative selection), and as people who inherited resistance mutations contributed more to the next generation (positive selection), TB gradually evolved from a severe, acute, systemic infection to a rare chronic lung infection. This was true until the late 1980s, when shifting events created conditions ideal for the infection’s return.

At first, complacency led to the resurgence of TB. Funding for TB research dried up because the infection was considered “cured.” Patients thought themselves cured when

antibiotics treated the symptoms in a few months, abandoning the drugs yet unknowingly continuing to spread the bacteria. As air travel became more commonplace, people began spreading different strains of the bacteria around the world. Treatment in the 1950s—isolating patients for 18 months or longer in facilities called sanatoria—was actually more effective by quarantining infectious individuals. Then AIDS happened.

AIDS shattered immunity, providing millions of vulnerable human lungs to support TB bacteria—which had never vanished, just retreated into milder forms. With so many more lungs to infect, bacterial mutations began to accumulate, and variants resistant to antibiotic drugs arose. Today, one-third of the 40 million HIV-infected people worldwide also have TB. Someone with HIV faces a fifty-fold increased risk of contracting TB. Each disease speeds the course of the other. In terms of public health, the most frightening aspect of the problem of dual infection is that a person with HIV can pass TB to anyone in just a sneeze or cough.

Further evidence of evolution is that the bacteria that cause TB are becoming resistant to many types of antibiotic drugs and the genetic diversity of the bacteria is increasing. Resistance genes are passed on genetic elements called transposons (discussed in section 11.4), which means that a bacterium can pick up several resistances at once. Today, multidrug-resistant (MDR) TB accounts for about 10 percent of cases in many nations, and up to 50 percent of cases in China, Russia, and India. In these three nations, poverty and faltering health care systems have introduced the stress and malnutrition that impair immunity, which allows TB to take hold. Symptoms of MDR TB are bloody cough, weight loss, lack of appetite, fever, and night sweats.

The HIV epidemic and stressful environments such as prisons and orphanages in Russia continue to provide breeding grounds for TB, but the problem may be coming under control, thanks to a program from the World Bank, World Health Organization, and the Global Fund. The program is introducing earlier diagnosis and treatment, monitoring of antibiotic treatment for 6 to 8 months, and improved reporting and surveillance of cases. However, the resurgence of TB should remind us never to underestimate the evolution that operates in all organisms—and does so unpredictably. **Reading 15.1** discusses antibiotic resistance.

Evolving HIV

Because the RNA or DNA of viruses replicates often and errors are not repaired, viral mutations accumulate rapidly. Like bacteria, the viruses in a human body form a population,

including naturally occurring genetic variants. In HIV infection, natural selection controls the diversity of HIV genetic variants within a human body as the disease progresses. The human immune system and drugs to slow the infection become the environmental factors that select (favor) resistant viral variants.

HIV infection can be divided into three stages, both from the human and the viral perspective (**figure 15.12**). A person infected with HIV may experience an initial acute phase, with symptoms of fever, night sweats, rash, and swollen glands. In a second period, lasting from 2 to 15 years, health usually returns. In a third stage, immunity collapses, the virus replicates explosively, and opportunistic infections and cancer eventually cause death.

The HIV population changes and expands throughout the course of infection, even when the patient seems to stay the same for a long time. New mutants continuously arise, and they alter such traits as speed of replication and the patterns of molecules on the viral surface.

In the first stage of HIV infection, as the person battles acute symptoms, viral variants that replicate swiftly predominate. In the second stage, the immune system starts to fight back and symptoms abate, as viral replication slows and many viruses are destroyed. Now natural selection acts, and certain viral variants reproduce and mutate, giving rise to a diverse viral population. Ironically, drugs used to treat AIDS may further select against the weakest HIV variants. Gradually, the

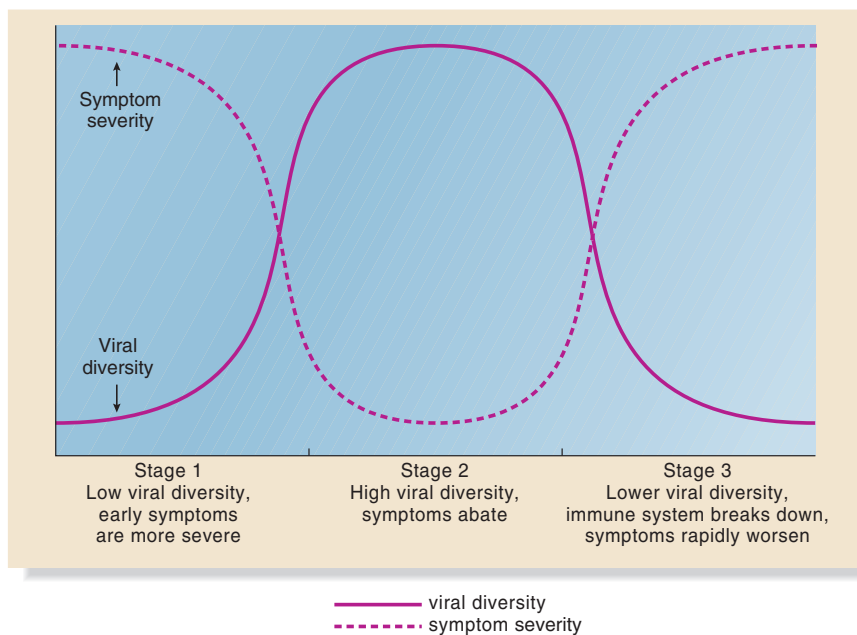


Figure 15.12 Natural selection of HIV. Natural selection controls the genetic diversity of an HIV population in a person's body. Before the immune system gathers strength, and after it breaks down, HIV diversity is low. A rapidly reproducing viral strain predominates, although new mutations continually arise. During the 2- to 15-year latency period, viral variants that can evade the immune system gradually accumulate.

Reading 15.1

Antibiotic Resistance: The Rise of MRSA

Many antibiotic drugs are no longer effective in treating bacterial infections. The reason is the interplay between mutation and natural selection.

Our bodies harbor populations of bacteria that have genetic variants, some of which enable the microorganisms to survive in the presence of a particular antibiotic drug. When a sick person takes the drug, symptoms abate as sensitive bacteria die. The resistant mutants reproduce, taking over the niche the antibiotic-sensitive bacteria vacated. Soon, the person has enough antibiotic-resistant bacteria to feel ill again. Usually antibiotic resistance genes already exist in the bacterial populations, and exposure to the drug selects the resistant bacteria. However, some antibiotics actually induce mutation.

Resistant bacteria circumvent antibiotic actions in several ways. Penicillin kills bacteria by tearing apart their cell walls. Resistant microbes produce enzyme variants that dismantle penicillin, or have altered cell walls that the drug cannot bind. Erythromycin, streptomycin, tetracycline, and gentamicin kill bacteria by attacking their ribosomes, which are different from ribosomes in a human. Drug-resistant bacteria have ribosomes that the drugs cannot bind.

Bacteria become resistant in two ways. Their DNA can mutate, passing the resistance from one bacterial generation to the next by cell division. Or, groups of resistance genes are passed on transposons, which move from cell to cell as part of DNA circles called plasmids. Bacteria usually pass transposons to similar bacteria, but in the unnatural environment of a hospital, genes may flit to any bacterium, and drug resistances are passed quickly. This is what has happened with infection by the bacterium *Staphylococcus aureus*.

S. aureus is normally present in low numbers in the nose and on the skin, but in high numbers it causes pimples and boils, food poisoning, toxic shock syndrome, pneumonia, and surgical wound infections (figure 1). *S. aureus* infection is particularly dangerous in hospitals, spreading rapidly among people unable to fight it. This common bacterium became resistant to penicillin soon after the drug was introduced in the 1940s. A related penicillin, methicillin, worked for a time, but resistant bacterial strains appeared suddenly in 2000, at such an alarming rate that the microorganism has its own acronym: MRSA, for methicillin-resistant *Staph aureus*. Doctors have turned to

another antibiotic, vancomycin, to treat MRSA, but the effort may be too late.

DNA sequencing revealed that in one hospital, *S. aureus* picked up vancomycin resistance from another type of bacterium. A foot ulcer in a dialysis patient in Detroit harbored vancomycin-resistant *Enterococcus faecalis* as well as two types of *S. aureus*, one resistant to the antibiotic and one sensitive. By sequencing the plasmids that included the resistance gene, investigators deduced that *S. aureus* picked up an *E. faecalis* plasmid bearing a vancomycin resistance gene called *vanA*. Then the *vanA* gene jumped to an *S. aureus* plasmid. Since then, similar scenarios of drug resistance gene sharing among microorganisms have happened in many countries.

The frightening part of the rise of MRSA infection relates back to natural selection, which in this case benefits the pathogen, and not us. That is, bacteria that can resist the drugs that we use to fight them will survive and reproduce, ensuring that *Staphylococcus aureus* infection continues.

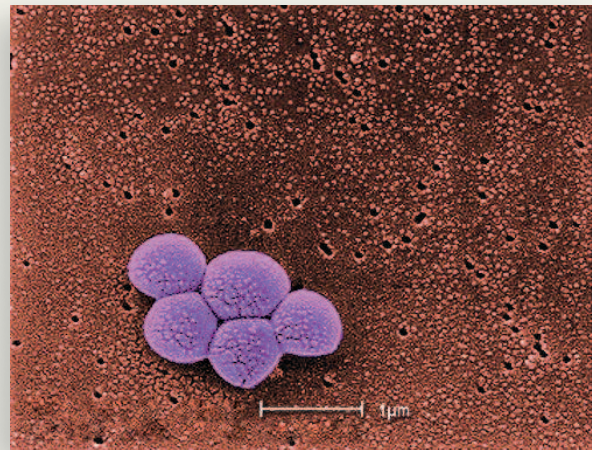


Figure 1 Antibiotic resistance. These *Staphylococcus aureus* bacteria are resistant to methicillin and several other antibiotics.

HIV population overtakes the immune system cells, but years may pass before immunity begins to noticeably decline. The third stage, full-blown AIDS, occurs when the virus overwhelms the immune system. With the selective pressure off, viral diversity again diminishes, and the fastest-replicating variants predominate. HIV wins.

The entire scenario of HIV infection reflects the value of genetic diversity—to enable the survival of a population or species in the face of an environmental threat. When that

threat—an immune system attack or drugs—wipes out sensitive variants, one genotype may prevail.

Knowing that HIV diversifies early in the course of infection has yielded clinical benefits. This is why patients now take combinations of drugs right after diagnosis. The drugs act in different ways to squelch several viral variants simultaneously, slowing the course of the infection. For many people, thanks to declining viral genetic diversity, HIV infection has become a chronic illness rather than the swift killer that it was when the epidemic began.

Balanced Polymorphism

If natural selection eliminates individuals with detrimental phenotypes from a population, then how do harmful mutant alleles remain in a gene pool? Harmful recessive alleles are replaced in two ways: by new mutation, and by persistence in heterozygotes.

Sometimes, a recessive condition remains particularly prevalent because the heterozygote enjoys a health advantage, such as being resistant to an infectious disease or able to survive an environmental threat. This “heterozygous advantage” that maintains a recessive, disease-causing allele in a population is called **balanced polymorphism**. Recall that *polymorphism* means variant; the effect is *balanced* because the protective effect of the noninherited condition counters the negative effect of the deleterious allele, maintaining its frequency in the population. Balanced polymorphism is a type of balancing selection, which more generally refers to maintaining heterozygotes in a population. A few examples follow, and these and others are summarized in **table 15.5**.

Sickle Cell Disease and Malaria

Sickle cell disease is an autosomal recessive disorder that causes anemia, joint pain, a swollen spleen, and frequent, severe infections. It is the classic example of balanced polymorphism: carriers are resistant to malaria, or develop very mild cases.

Malaria is an infection by the parasite *Plasmodium falciparum* and related species that causes debilitating cycles of chills and fever. The parasite spends the first stage of its life cycle in the salivary glands of the mosquito *Anopheles gambiae*. When an infected female mosquito draws blood from a human, malaria parasites enter red blood cells, which transport

the parasites to the liver. The red blood cells burst, releasing parasites throughout the body.

In sickle cell disease, many red blood cells burst prematurely, which expels the parasites before they can cause rampant infection. The blood of a person with sickle cell disease is also thicker than normal, which may hamper the parasite’s ability to infect. A sickle cell disease carrier’s blood has enough abnormal cells to be inhospitable to the malaria parasite—but usually not enough to cause the blocked circulation of sickle cell disease.

A clue to the protective effect of being a carrier for sickle cell disease came from striking differences in the incidence of the two diseases in different parts of the world (**figure 15.13**). In the United States, 8 percent of African Americans are sickle cell carriers, whereas in parts of Africa, up to 45 to 50 percent are carriers. Although Africans had known about a painful disease that shortened life, the sickled cells weren’t recognized until 1910 (see section 12.2). In 1949, British geneticist Anthony Allison found that the frequency of sickle cell carriers in tropical Africa was higher in regions where malaria rages all year long. Blood tests from children hospitalized with malaria showed that nearly all were homozygous for the wild type sickle cell allele. The few sickle cell carriers among them had the mildest cases of malaria. Was malaria enabling the sickle cell allele to persist by felling people who did not inherit it? The fact that sickle cell disease is rarer where malaria is rare supports the idea that sickle cell heterozygosity protects against the infection.

Further evidence of a sickle cell carrier’s advantage in a malaria-ridden environment is the fact that the rise of sickle cell disease parallels the cultivation of crops that provide breeding grounds for *Anopheles* mosquitoes. About 1000 B.C., Malayo-Polynesian sailors from southeast Asia traveled in

Table 15.5 Balanced Polymorphism

Disease 1 (inherited, carrier)	Protects against →	Disease 2	Because →	Mechanism	References
Sickle cell disease		Malaria		Abnormal red blood cells cannot retain parasites	Section 12.2
G6PD deficiency		Malaria		Parasite cannot reproduce in abnormal red blood cells	Section 12.5
PKU		Fungal infection in fetuses		Elevated phenylalanine inactivates fungal toxin	Sections 5.2, 14.1, 15.6
Prion protein mutation		Transmissible spongiform encephalopathy		Prion protein cannot misfold in presence of infectious prion protein	Figure 10.22, Section 12.5, Reading 10.1
CF		Diarrheal disease (cholera, typhus)		Fewer chloride channels in intestinal cells prevent water loss	Sections 14.1, 14.3, Reading 2.2
Smith-Lemli-Opitz syndrome		Cardiovascular disease		Lowered serum cholesterol	MIM 270400 (multiple birth defects, mental retardation)

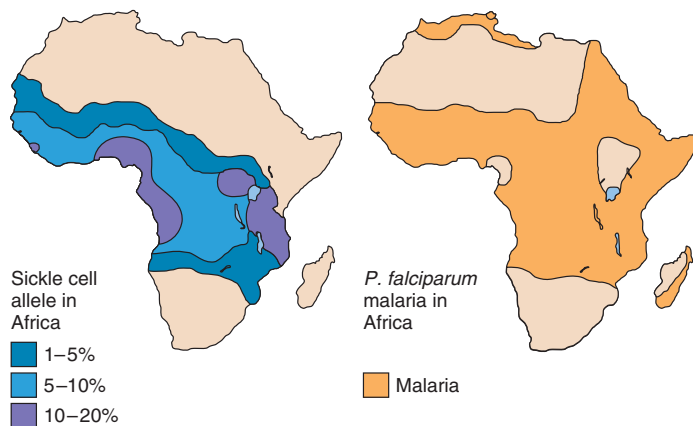


Figure 15.13 **Balanced polymorphism.** Comparing the distribution of people with malaria and people with sickle cell disease in Africa reveals balanced polymorphism. Carriers for sickle cell disease are resistant to malaria because changes in the blood caused by the sickle cell allele are not severe enough to impair health, but they do inhibit the malaria parasite.

canoes to East Africa, bringing new crops of bananas, yams, taros, and coconuts. When the jungle was cleared to grow these crops, the open space provided breeding grounds for the mosquitoes. The insects, in turn, offered a habitat for part of the life cycle of the malaria parasite.

The sickle cell allele may have been brought to Africa by people migrating from Southern Arabia and India, or it may have arisen directly by mutation in East Africa. However it happened, people who inherited one copy of the sickle cell allele survived or never contracted malaria—the essence of natural selection. These carriers had more children and passed the protective allele to approximately half of them. Gradually, the frequency of the sickle cell allele in East Africa rose from 0.1 percent to 45 percent in 35 generations. Carriers paid the price for this genetic protection, however, whenever two of them produced a child with sickle cell disease.

A cycle set in. Settlements with large numbers of sickle cell carriers escaped debilitating malaria. They were strong enough to clear even more land to grow food—and support the disease-bearing mosquitoes.

Prion Disease and Cannibalism

Being a heterozygote for the prion protein gene may protect against the disorders of protein folding called transmissible spongiform encephalopathies (see figure 10.21, Reading 10.1, and section 12.5). The best studied such illness is kuru, which caused brain degeneration among the Foré people in Papua New Guinea until the Australian government halted the practice of ritual cannibalism in the mid-1950s. A recent investigation of the prion protein gene among 30 elderly Foré women who had eaten brains revealed 23 heterozygotes. Only 15 were predicted based on Hardy-Weinberg equilibrium observed among 140 Foré who had not eaten brains. In the heterozygotes, some of the normal prion proteins have a valine at amino acid position 129, and some a methionine. The different amino acids

somehow prevent infectious misfolding in the presence of abnormal prion protein—as happens in cannibalism. All of the people in the United Kingdom who have developed variant CJD, the human form of “mad cow disease,” have only methionine at position 129.

The overrepresentation of heterozygotes among the Foré survivors led to the hypothesis that balancing selection has favored this genotype in the population, and that cannibalism may have been the driving force. That is, homozygotes who were cannibals died of a prion disorder before reproducing, leaving the resistant heterozygotes to slowly accumulate in the population.

This new genetic view of cannibalism supports anthropological evidence that eating human flesh has occurred in many times and places, from Neanderthal caves in France and Croatia to the American Southwest. Evidence of cannibalism includes human bones damaged in ways similar to the bones of animals prepared for consumption, such as scratch marks to remove muscle and signs of breaking and crushing to obtain marrow. Biochemical evidence for past cannibalism includes human myoglobin, found only in human muscle, in fossilized human excrement.

Cystic Fibrosis and Diarrheal Disease

Balanced polymorphism may explain why CF is so common—its cellular defect protects against diarrheal illnesses such as cholera and typhus. Diarrheal disease epidemics have left their mark on many human populations, and continue to be a major killer in the developing world.

Severe diarrhea rapidly dehydrates the body and leads to shock, kidney and heart failure, and death in days. In cholera, bacteria produce a toxin that opens chloride channels in cells of the small intestine. As salt (NaCl) leaves the intestinal cells, water rushes out, producing diarrhea. The CFTR protein does just the opposite, closing chloride channels and trapping salt and water in cells, which dries out mucus and other secretions. A person with CF is very unlikely to contract cholera, because the toxin cannot open the chloride channels in the small intestine cells.

CF carriers enjoy the mixed blessing of balanced polymorphism. They do not have enough abnormal chloride channels to cause the labored breathing and clogged pancreas of CF, but they have enough of a defect to block the cholera toxin. During the devastating cholera epidemics that have occurred throughout history, individuals carrying mutant CF alleles had a selective advantage, and they disproportionately transmitted those alleles to future generations. Cholera epidemics still happen.

Because CF arose in western Europe and cholera originated in Africa, an initial increase in CF heterozygosity may have been a response to a different diarrheal infection—typhoid fever. The causative bacterium, *Salmonella typhi*, rather than producing a toxin, enters cells lining the small intestine—but only if CFTR channels are present. The cells of people with severe CF manufacture CFTR proteins that never reach the cell

surface, and therefore no bacteria get in. Cells of CF carriers admit some bacteria. Protection against infections that produce diarrhea may therefore have kept CF in populations.

Key Concepts

1. Different alleles are more likely to confer a survival advantage in different environments. Cycles of infectious disease prevalence and virulence often reflect natural selection.
2. In balanced polymorphism, a disease-causing allele persists because heterozygotes resist a certain infectious illness or environmental condition.

15.6 Putting It All Together: PKU

Monitoring allele frequencies in populations can reveal the effects of nonrandom mating, migration, genetic drift, mutation, and natural selection on evolution. **Figure 15.14** summarizes the forces acting alone, and **table 15.6** lists the examples in the chapter with the mechanisms that they illustrate. Often, historical, archeological, and linguistic evidence can help us to understand the complex interactions of these forces. Consider, again, PKU, which causes mental retardation unless a specific diet is followed from birth.

The diversity of PKU mutations suggests that the disease has arisen more than once. Mutations common to many groups of people probably represent more ancient mutations that occurred before groups spread and separated. In contrast, mutations found only in a small geographical region are more likely to be of recent origin, perhaps kept apart by genetic drift. These mutations have had less time to spread. For example, Turks, Norwegians, French Canadians, and Yemeni Jews have their own PKU alleles. Analysis of the frequencies of PKU mutations in different populations, plus logic, can reveal the roles that genetic drift, mutation, and balanced polymorphism have played in maintaining the mutation.

A high mutation rate cannot be the sole reason for the continued prevalence of PKU because some countries continue to have only one or two mutations. If the gene mutated frequently, all populations would have several different types of PKU mutations. This is not so.

In some isolated populations, migration and the founder effect have maintained certain PKU alleles. In most populations, point mutations in the phenylalanine hydroxylase (*PAH*) gene cause PKU. However, all Yemeni Jews in Israel who have PKU have a large deletion. This mutation spread from northern Africa to Israel. To track it, researchers tested for the telltale deletion in the grandparents of the 22 modern Yemeni Jewish families with PKU in Israel. The earliest court and religious records identify two families with PKU in San'a, the capital of Yemen. By 1809, religious persecution and hard economic times led nine families carrying the mutation to migrate north and settle in three towns, then spread farther (**figure 15.15**).

PKU may have persisted because being a heterozygote protects against certain fungal infections spread in tainted grains, particularly in Ireland and Scotland, where the inherited disease is the most prevalent. PKU carriers have elevated phenylalanine levels in their blood that are not enough to cause PKU symptoms, but high enough to inactivate a fungal

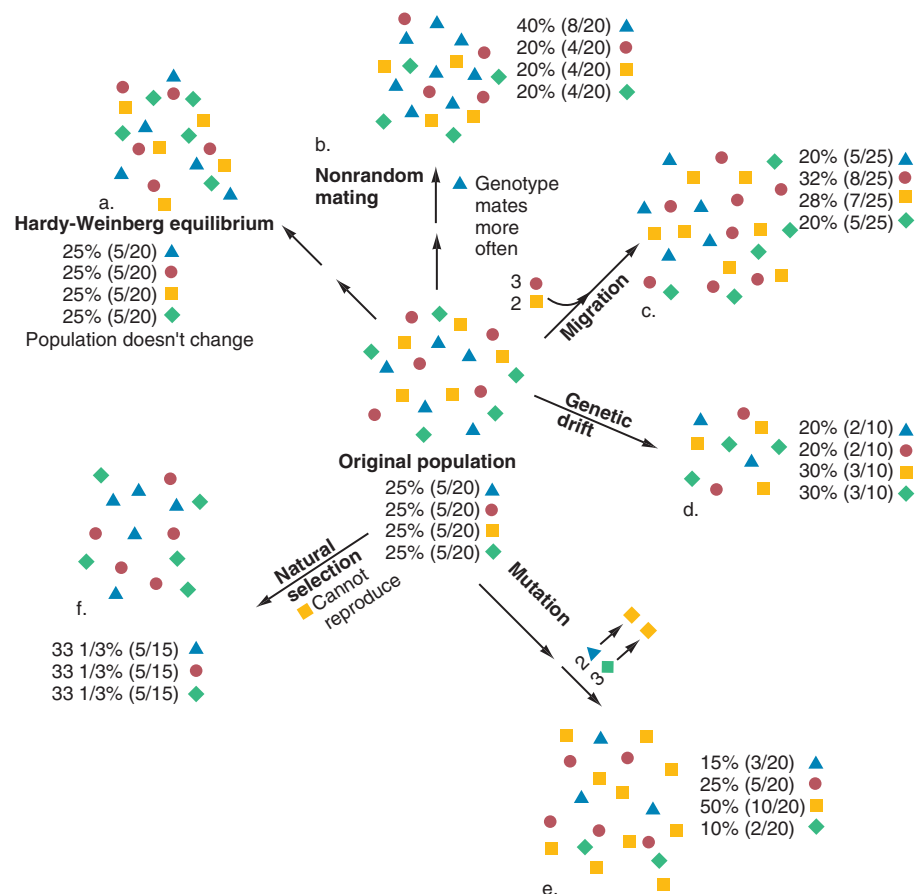


Figure 15.14 Forces that change allele frequencies.

Table 15.6 Forces that Change Allele Frequencies

Mechanism of Allele Frequency Change	Examples
Nonrandom mating	Agriculture Cape population and Arnold Hopi Indians with albinism Genghis Khan's Y chromosome
Migration	Consanguinity Galactokinase deficiency in Europe ABO blood type distribution Clines along the Nile and in Italy
Genetic drift	
Founder effect	Fumarate deficiency in Arizona/Utah Disorders among French-Canadians, Dunkers, Old Order Amish and Mennonites Afrikaners and porphyria variegata
Population bottleneck	Pingelapese blindness Cheetahs Chmielnicki massacre
Mutation	Chapters 12 and 13
Natural selection	Lactose intolerance TB incidence and virulence HIV infection Antibiotic resistance Sickle cell disease and malaria Prion disease and cannibalism CF and diarrheal disease

poison, ochratoxin A, that harms fetuses. During the famines that have periodically plagued this part of the world, starving people ate moldy grain. If PKU carriers eating the grain were more likely to have healthy children, the mutant allele would have increased in the population, thanks to balanced polymorphism. The Vikings spread PKU eastward.

Key Concepts

1. PKU originated more than once.
2. Genetic drift, balanced polymorphism, and perhaps mutation have affected its prevalence.

15.7 Eugenics

We usually think of artificial selection in the context of Darwin's pet pigeons or purebred cats, dogs, and horses. We also practice artificial selection through control of our own reproduction, through individual choices as well as at the societal level.

Some people attempt to control the genes in their offspring. They do this by seeking mates with certain characteristics, by choosing egg or sperm donors with particular traits, or by ending pregnancies after a test reveals a devastating disorder. On a societal level, **eugenics** refers to programs that control human reproduction with the intent of changing the genetic structure of the population. Eugenics works in two directions. Positive eugenics creates incentives for reproduction among those considered superior; negative eugenics interferes with reproduction of those judged inferior. Obviously, eugenic measures are highly subjective. **Table 15.7** lists some famous examples of eugenics, and *Bioethics: Choices for the Future* (pg. 301) considers a highly personal viewpoint.

The word "eugenics" was coined in 1883 by Sir Francis Galton to mean "good in birth." He defined eugenics as "the science of improvement of the human race germplasm through better breeding." One vocal supporter of the eugenics movement was Sir Ronald Aylmer Fisher. In 1930, he tried to apply the principles of population genetics to human society, writing that those at the top of a society tend to be "genetically infertile," producing fewer children than the less-affluent classes. This, he claimed, was the reason why civilizations ultimately topple. He offered several practical suggestions to remedy this, including state monetary gifts to high-income families for each child born to them.

Early in the twentieth century, eugenics focused on maintaining purity. One prominent geneticist, Luther Burbank, realized the value of genetic diversity at the beginning of a eugenic effort. Known for selecting interesting plants and crossing them to breed plants with useful characteristics, Burbank in 1906 applied his ideas to people. In a book called *The Training of the Human Plant*, he encouraged immigration to the United States so that advantageous combinations of traits would appear as the new Americans interbred. Burbank's plan ran into problems,

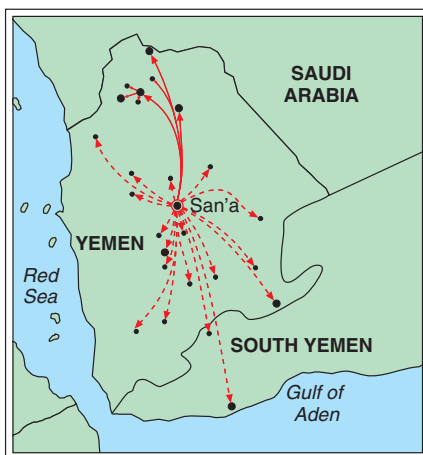


Figure 15.15 The origin of PKU. The deletion in Israeli Yemeni Jews probably arose in San'a, Yemen, in the mid-eighteenth century. The allele spread northward as families moved from San'a in 1809 (solid arrows) and subsequently spread to other regions (broken arrows), and eventually Israel.

Source: Data from Smadar Avigad, et al., *A single origin of phenylketonuria in Yemenite Jews*, *Nature* 344:170, March 8, 1990.

Table 15.7 A Chronology of Eugenics-Related Events

1883	Sir Francis Galton coins the term <i>eugenics</i> .
1889	Sir Francis Galton's writings are published in the book <i>Natural Inheritance</i> .
1896	Connecticut enacts law forbidding sex with a person who has epilepsy or is "feeble-minded" or an "imbecile."
1904	Galton establishes the Eugenics Record Office at the University of London to keep family records.
1907	First eugenic law in the United States orders sterilization of institutionalized mentally retarded males and criminal males when experts recommend it.
1910	Eugenics Record Office founded in Cold Spring Harbor, New York, to collect family and institutional data.
1924	Immigration Act limits entry into the United States of "idiots, imbeciles, feeble-minded, epileptics, insane persons," and restricts immigration to 7 percent of the U.S. population from a particular country according to the 1890 census—keeping out those from southern and eastern Europe.
1927	Supreme Court (<i>Buck vs. Bell</i>) upholds compulsory sterilization of the mentally retarded by a vote of 8 to 1, leading to many state laws.
1934	Eugenic sterilization law of Nazi Germany orders sterilization of individuals with conditions thought to be inherited, including epilepsy, schizophrenia, and blindness, depending upon rulings in Genetic Health Courts.
1939	Nazis begin killing 5,000 children with birth defects or mental retardation, then 70,000 "unfit" adults.
1956	U.S. state eugenic sterilization laws are repealed, but 58,000 people have already been sterilized.
1965	U.S. immigration laws reformed, lifting many restrictions.
1980s	California's Center for Germinal Choice established, where Nobel Prize winners can deposit sperm to inseminate selected women.
1990s	In the U.S., state laws passed to prevent health insurance or employment discrimination based on genotype.
2003	Many governments recommend certain genetic tests, and have legislation to prevent genetic discrimination. In the U.S., protective legislation is still in discussion.
2004	Genocide of black Africans occurs in Sudan.
2008	Federal genetic anti-discrimination legislation finalized in U.S.
2009	Genetic information nondiscrimination act enacted.

however, at the selection stage, which allowed only those with "desirable" traits to reproduce.

On the East Coast of the United States, Charles Davenport led the eugenics movement. In 1910, he established the Eugenics Record Office at Cold Spring Harbor, New York. There he headed a massive effort to compile data from institutions, prisons, circuses, and general society. He attributed nearly every trait to a single gene. "Feeble-mindedness," for example, was an autosomal recessive trait. It was a catch-all phrase for a person with low intelligence (as measured on an IQ test) and such abnormalities as "criminality," "promiscuity," and "social dependency." In one famous case, a young woman named Carrie Buck was ordered to be sterilized when she, her mother, and her illegitimate infant daughter Vivian were declared feeble-minded. Carrie had been raped by a relative of her foster parents, and was actually an average student. **Figure 15.16** shows the pedigree for Carrie Buck and her "inherited trait" of feeble-mindedness.

Other nations practiced eugenics. From 1934 until 1976, the Swedish government forced certain individuals to be sterilized as part of a "scientific and modern way of changing society

for the better," according to one historian. At first, only mentally ill people were sterilized, but poor, single mothers were later included. The women's movement in the 1970s pushed for an end to forced sterilizations.

In 1994, China passed the Maternal and Infant Health Care Law, which proposes "ensuring the quality of the newborn population" and forbids procreation between two people if physical exams show "genetic disease of a serious nature . . . that may totally or partially deprive the victim of the ability to live independently, that [is] highly possible to recur in generations to come, and that [is] medically considered inappropriate for reproduction." Such "genetic diseases" include mental retardation, mental illness, and seizures, conditions that are ill-defined in the law and are not necessarily inherited.

Another guise of eugenics is war, if the fighting groups differ genetically. Throughout history, war and conflict have altered gene pools, sometimes dramatically. These effects are eugenic when they take the form of rape of women of one group by men from another, with the intent of "diluting" the genes of the rape victims. During the Rwandan genocide of 1994,

Bioethics: Choices for the Future

Two Views of Neural Tube Defects

Genetic tests enable people to make reproductive choices that can alter allele frequencies in populations. Identifying carriers of a recessive illness, who then may decide not to have children together, is one way to remove disease-causing alleles from a population. Screening pregnant women for fetal anomalies, then terminating affected pregnancies, also alters disease prevalence and, if the disorder has a genetic component, allele frequencies. This is the case for neural tube defects (NTDs), which are multifactorial.

An NTD forms at the end of the first month, when the embryo's neural tube does not completely close. An opening in the head (anencephaly) usually ends in miscarriage, stillbirth, or a newborn who dies within days. An opening in the spinal cord (spina bifida) causes paralysis but the person can live into adulthood and have normal intelligence. Surgery can help.

In 1992, the Centers for Disease Control and Prevention concluded that taking the vitamin folic acid in pregnancy lowers the risk of NTD recurrence from 3 to 4 percent to 1.5 to 2 percent. Women who had had an affected child began taking large doses before conception. But when epidemiologists tried to monitor how well it was working, they faced a problem—prevalence of NTDs was greatly underestimated. This happened because the statistics on NTD prevalence—vital to discovering whether folic acid was preventing the defect—included only newborns, stillborns, and older fetuses. Most reports did not account for pregnancies terminated following prenatal diagnosis. These pregnancies caused the underreporting of anencephaly by 60 to 70 percent, and of spina bifida, by 20 to 30 percent in some states.

A Personal View

Blaine Deatherage-Newsom has a different view of population screening for NTDs because he has one (see photo). Blaine was born in 1979 with spina bifida. Paralyzed from the armpits down, he has endured much physical pain, but he has also achieved a great deal. While in high school, he put the question, "If we had the technology to eliminate disabilities from the population, would that be good public policy?" on the Internet—initiating a global discussion. He wrote:

I was born with spina bifida and hydrocephalus. I hear that when parents have a test and find out that their unborn child has spina bifida, in more than 95 percent of the cases they choose to have an abortion. I also went to an exhibit at the Oregon Museum of Science and Industry several years ago where the exhibit described a child born with spina bifida and hydrocephalus, and . . . asked people to vote on whether the child should live or die. I voted that the child should live, but when I voted, the child was losing by quite a few votes.

When these things happen, I get worried. I wonder if people are saying that they think the world would be a better place without me. I wonder if people just think the lives of people with disabilities are so full of misery and suffering that they think we would be better off dead. It's true that my life has suffering (especially when I'm having one of my 11 surgeries so far), but most

of the time I am very happy and I like my life very much. My mom says she can't imagine the world without me, and she is convinced that everyone who has a chance to know me thinks that the world is a far better place because I'm in it.

Today Blaine works for a not-for-profit organization that refurbishes computer equipment for community service organizations.

Questions for Discussion

1. Is the decision to end a pregnancy that would otherwise lead to the birth of a child with a neural tube defect a eugenic measure or not? State a reason for your answer.
2. People with certain medical conditions or limitations, such as those with hearing loss, object to genetic tests that would ultimately decrease their numbers in the population. How would you feel if you had such a condition?
3. Do you think that eugenics will resurge as personal genome sequencing and genetic testing become more widespread?

Excerpt by Blaine Deatherage-Newsom, "If we could eliminate disabilities from the population, should we? Results of a survey on the Internet." Reprinted by permission.



Blaine Deatherage-Newsom as a teen.

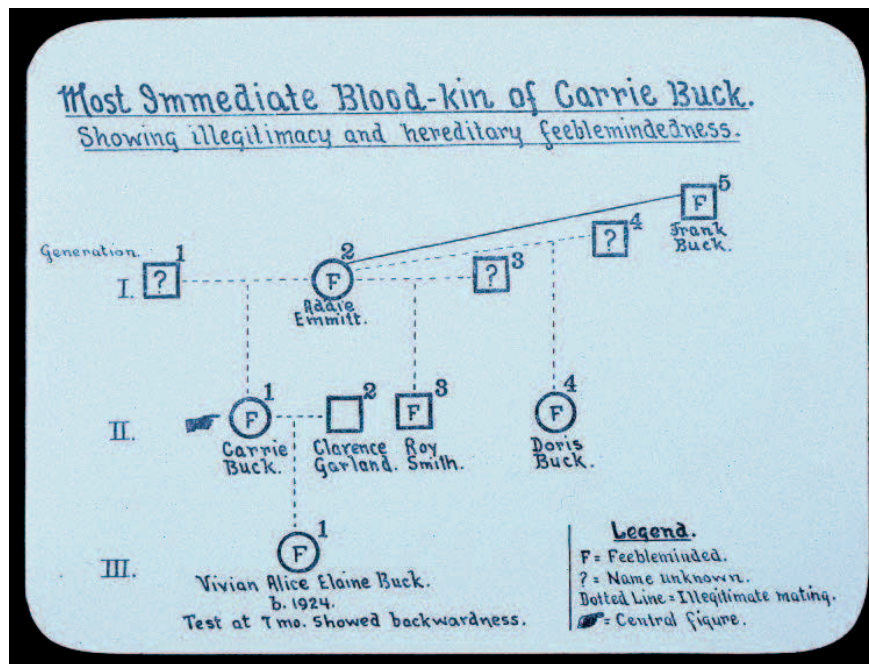


Figure 15.16 Eugenics sought to abolish “feeble-mindedness.” In 1927, 17-year-old Carrie Buck, of Charlottesville, stood trial for the crime of having a mother who lived in an asylum for the feeble-minded, and having a daughter out-of-wedlock (following rape) also deemed feeble-minded, as was Carrie herself, though she was a B student in school. Ruled Sir Oliver Wendell Holmes, Jr., “three generations of imbeciles are enough.” Carrie Buck made history as the first person to be sterilized to prevent having another “socially inadequate offspring.”

for example, Hutu policy was to rape Tutsi women—250,000 of them. Genocide by rape has been ongoing in the eastern Democratic Republic of the Congo since 1996. Since 2003 in Darfur, part of western Sudan in Africa, Arab militia have systematically attacked black Africans, killing men and children and repeatedly raping women. In Rwanda, Congo, and Darfur, the conquerors have claimed that their intent is to diminish the genetic contributions of their victims and spread their own genes. That’s eugenics.

Because genetic technologies may affect reproductive choices and can influence which alleles are passed to the next generation, modern genetics has been compared to eugenics. Medical genetics and eugenics differ in their overall goals. Eugenics aims to allow only people with certain “valuable” genotypes to reproduce, for the supposed benefit of the population as a whole. The goal of medical genetics, in contrast, is usually to prevent suffering on a family level.

Humanity arose in Africa some 200,000 years ago. As pockets of peoples spread across the globe, our behaviors and intelligence introduced culture. The next chapter explores some of our journeys, through clues to the past in the sequences of our DNA.

Key Concepts

1. Eugenics is the control of individual human reproduction for societal goals, maximizing the genetic contribution of those deemed acceptable (positive eugenics) and minimizing the contribution from those considered unacceptable (negative eugenics).
2. Some people consider modern genetic screening practices eugenic, but genetic testing usually aims to prevent or alleviate human suffering.
3. Wars may have eugenic consequences.

Summary

15.1 Nonrandom Mating

1. Hardy-Weinberg equilibrium assumes all individuals mate with the same frequency and choose mates without regard to phenotype. This rarely happens. We choose mates based on certain characteristics, and some people have many more children than others.

2. DNA sequences that do not cause a phenotype important in mate selection or reproduction may be in Hardy-Weinberg equilibrium.
3. Consanguinity increases the proportion of homozygotes in a population, which may lead to increased incidence of recessive illnesses or traits.

15.2 Migration

4. **Clines** are changes in allele frequencies from one area to another.
5. Clines may reflect geographical barriers or linguistic differences and may be either abrupt or gradual.
6. Human migration patterns through history explain many cline boundaries. Forces behind migration include escape from persecution and a nomadic lifestyle.

15.3 Genetic Drift

7. **Genetic drift** occurs when a small population separates from a larger one, or its members breed only among themselves, perpetuating allele frequencies not characteristic of the larger population due to chance sampling.
8. A **founder effect** occurs when a few individuals found a settlement and their alleles form a new gene pool, amplifying their alleles and eliminating others.
9. A **population bottleneck** is a narrowing of genetic diversity that occurs after many members of a population die and the few survivors rebuild the gene pool.

15.4 Mutation

10. Mutation continually introduces new alleles into populations. It occurs as a consequence of DNA replication errors.
11. Mutation does not have as great an influence on disrupting Hardy-Weinberg equilibrium as the other factors.
12. The **genetic load** is the collection of deleterious alleles in a population.

15.5 Natural Selection

13. Environmental conditions influence allele frequencies via **natural selection**. Alleles that do not enable an individual to reproduce in a particular environment are selected against and diminish in the population, unless conditions change. Beneficial alleles are retained.
14. In **balanced polymorphism**, the frequencies of some deleterious alleles are maintained when heterozygotes have a reproductive advantage under certain conditions.

15.6 Putting It All Together: PKU

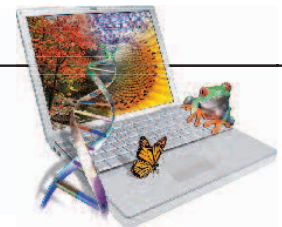
15. Frequencies of different mutations in different populations provide information on the natural history of alleles and on the relative importance of nonrandom mating, genetic drift, and natural selection in deviations from Hardy-Weinberg equilibrium.

15.7 Eugenics

16. **Eugenics** is the control of individual reproduction to serve a societal goal.
17. Positive eugenics encourages those deemed acceptable or superior to reproduce. Negative eugenics restricts reproduction of those considered inferior. Eugenics extends the concept of natural selection and Mendel's laws but does not translate well into practice.
18. Some aspects of genetic technology affect reproductive choices and allele frequencies, but the goal is to alleviate or prevent suffering, not to change society.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

1. Give examples of how each of the following can alter allele frequencies from Hardy-Weinberg equilibrium:
 - a. nonrandom mating
 - b. migration
 - c. a population bottleneck
 - d. mutation
2. Explain the influence of natural selection on
 - a. the virulence of tuberculosis.
 - b. bacterial resistance to antibiotics.
 - c. the changing degree of genetic diversity in an HIV population during infection.
3. Why can increasing homozygosity in a population be detrimental?
4. How might a mutant allele that causes an inherited illness in homozygotes persist in a population?
5. Give an example of an inherited disease allele that protects against an infectious illness.
6. Explain how table 15.2 indicates that genetic drift has occurred among the Dunkers.
7. How does a founder effect differ from a population bottleneck?
8. Describe two scenarios in human populations, one of which accounts for a gradual cline, and one for an abrupt cline.
9. How do genetic drift, nonrandom mating, and natural selection interact?
10. Define:
 - a. founder effect
 - b. balanced polymorphism
 - c. genetic load

- How does a knowledge of history, sociology, and anthropology help geneticists to interpret allele frequency data?
- Cite three examples of eugenic actions or policies.
- Distinguish between positive and negative selection, and between positive and negative eugenics. How do selection and eugenics differ?

Applied Questions

- Begin with the original population represented at the center of Figure 15.14, and deduce the overall, final effect of the following changes:
 - Two yellow square individuals join the population when they stop by on a trip and stay awhile.
 - Four red circle individuals are asked to leave as punishment for criminal behavior.
 - A blue triangle man has sex with many females, adding five blue triangles to the next generation.
 - A green diamond female produces an oocyte with a mutation that adds a yellow square to the next generation.
 - A new infectious disease affects only blue triangles and yellow squares, removing two of each from the next generation.
- Before 1500 A.D., medieval Gaelic society in Ireland isolated itself from the rest of Europe, physically as well as culturally. Men in the group are called “descendants of Niall,” and they all share a Y chromosome inherited from a single shared ancestor. In the society, men took several partners, and sons born out of wedlock were fully accepted. Today, in a corner of northwest Ireland, one in five men has the “descendant of Niall” Y chromosome. In all of Ireland, the percentage of Y chromosomes with the Niall signature is 8.2 percent. In western Scotland, where the Celtic language is similar to Gaelic, 7.3 percent of the males have the telltale Niall Y. In the U.S., among those of European descent, it is 2 percent. Worldwide, the Niall Y chromosome makes up only 0.13 percent of the total. What concept from the chapter do the data illustrate?
- Fred Schnee, who teaches human genetics at Loras College in Iowa, offers a good example of genetic drift: seven castaways are shipwrecked on an island. The first mate has blue eyes, the others brown. A coconut falls on the first mate, killing him. The coconut accident is a chance event affecting a small population. Explain how this event would affect allele frequency, and offer another example of genetic drift.
- The Old Order Amish of Lancaster, Pennsylvania have more cases of polydactyly (extra fingers and toes) than the rest of the world combined. All of the affected individuals descend from the same person, in whom the dominant mutation originated. Does this illustrate a population bottleneck, a founder effect, or natural selection? Give a reason for your answer.
- Predict how natural selection might affect the frequency of alleles that protect against HIV infection in Africa a century from now, based on what you know about TB.
- The *MDR1* gene encodes the protein portion of a glycoprotein (called P-glycoprotein) that dots the surfaces of intestinal lining cells and T lymphocytes. An *MDR1* allele that is overexpressed became prevalent in West Africans because the encoded protein enables cells to pump out toxins, such as those produced when bacteria contaminate food. Although these people rarely have stomachaches, HIV drugs have little effect on them. If HIV infection continues to spread in West Africa, but food poisoning becomes less common as people learn to fully cook their food, what effect will natural selection have on this gene?
- The ability to taste bitter substances is advantageous in avoiding poisons, but might keep people from eating bitter vegetables that contain chemicals that protect against cancer. Devise an experiment, perhaps based on population data, to test either hypothesis—that the ability to taste bitter substances is either protective or harmful.
- Define microevolution. Give three examples, either from the chapter or from the news, that show microevolution going on right now.
- A mutation that removes the receptor for HIV on human cells also blocks infection by the bacterium that causes plague. Seven centuries ago, in Europe, the “Black Death” plague epidemic increased the protective allele in the population. Today it makes 3 million people in the United States and the United Kingdom resistant to HIV infection. Is the increase in incidence of this allele due to nonrandom mating or natural selection?
- Use the information in chapters 14 and 15 to explain why
 - porphyria variegata is more prevalent among Afrikaners than other South African populations.
 - many people among the Cape population in South Africa lose their teeth before age 20.
 - cystic fibrosis and sickle cell disease remain common.
 - the Pima Indians have an extremely high incidence of type 2 diabetes.
 - the Amish in Lancaster County and certain Pakistani groups have a high incidence of genetic diseases that are very rare elsewhere.
 - the frequency of the allele that causes galactokinase deficiency varies across Europe.
 - mitochondrial DNA sequences vary gradually in populations along the Nile River valley.
 - disease-causing *BRCA1* alleles are different in Jewish people of eastern European descent and African Americans.
- Which principles discussed in this chapter do the following classic science fiction plots illustrate?
 - In *When Worlds Collide*, the Earth is about to be destroyed, and 100 people are selected to colonize a new planet.
 - In *The Time Machine*, set in the distant future on Earth, one group of people is forced to live on the planet’s surface while another group is forced to live in caves. Over many

- years, they come to look and behave differently. The Morlocks that live below ground have dark skin, dark hair, and are very aggressive, whereas the Eloi that live above ground are blond, fair-skinned, and meek.
- c. In *Children of the Damned*, all of the women in a small town are suddenly made pregnant by genetically identical beings from another planet.
- d. In *The War of the Worlds*, Martians cannot survive on Earth because they are vulnerable to infection by terrestrial microbes.
- e. In Dean Koontz's novel *The Taking*, giant mutant fungi kill nearly everyone on Earth, sparing only young children and the few adults who protect them. The human race must re-establish itself from the survivors.
12. Ashkenazim, French Canadians, and people who live in southwestern Louisiana have a higher incidence of Tay-Sachs disease than other populations.
- a. Each of these groups has a different mutation. How is this possible?
- b. A controversial hypothesis proposes that the high incidence of Tay-Sachs disease and other genetic disorders that harm brain cells among the Ashkenazim reflects balanced polymorphism. Because brain cells are affected, carriers are, for reasons unknown, more intelligent and therefore had a survival advantage during periods of persecution because they could better use their wits to escape violence. What evidence might support the hypothesis?
13. Syndrome X consists of obesity, type 2 diabetes, hypertension, and heart disease. Researchers surveyed and sampled blood from nearly all of the 2,188 residents of the Pacific Island of Kosrae, and found that 1,709 of them are part of the same pedigree. The incidence of all of the symptoms of syndrome X is much higher in this population than for other populations. Suggest a reason for this finding, and indicate why it would be difficult to study these particular traits, even in an isolated population.
14. By which mechanisms discussed in this chapter do the following situations alter Hardy-Weinberg equilibrium?
- a. Ovalocytosis (MIM 166910) is caused by a beneficial mutation. A protein that anchors the red blood cell plasma membrane to the cytoplasm is abnormal, making the membrane so rigid that parasites that cause malaria cannot enter.
- b. In the mid-1700s, a multitoed male cat from England crossed the sea and settled in Boston, where he left behind quite a legacy of kittens—about half of whom also had extra digits. People loved the odd felines and bred them. Today, in Boston and nearby regions, multitoed cats are far more common than in other parts of the United States.
- c. Many slaves in the United States arrived in groups from Nigeria, which is an area in Africa with many ethnic subgroups. They landed at a few sites and settled on widely dispersed plantations. Once emancipated, former slaves in the South were free to travel and disperse.
- d. About 300,000 people in the United States have Alzheimer disease caused by a mutation in the presenilin-2 gene. They all belong to five families that came from two small villages in Germany that migrated to Russia in the 1760s and then to the U.S. from 1870 through 1920.
15. A challenging environment can either kill individuals whose genetic susceptibilities and characteristics make it difficult to survive or reproduce, or encourage such individuals to migrate to more comfortable surroundings. Describe the different effects of these alternatives on the genetic structure of the population.

Web Activities

16. Go to the Centers for Disease Control and Prevention website and access the journal *Emerging Infectious Diseases*. Using this resource, describe an infectious disease that is evolving, and cite the evidence for this.
17. Do a Google search for a pair of disorders listed in table 15.5 (balanced polymorphism) and discuss how the carrier status of the inherited disease protects against the second condition.
18. Go to the Image Archive on the American Eugenics Movement website. Look at several images, and either find one that presents a genetic disorder and describe it, or find an image that presents biologically incorrect information, and explain the error.

Case Studies and Research Results

19. The human population of India is divided into many castes, and the people follow strict rules governing who can marry whom. Researchers compared several genes among 265 Indians of different castes and 750 people from Africa, Europe, and Asia. The study found that the genes of higher Indian castes most closely resembled those of Europeans, and that the genes of the lowest castes most closely resembled those of Asians. In addition, maternally inherited genes (mitochondrial DNA) more closely resembled Asian versions of those genes, but paternally inherited genes (on the Y chromosome) more closely resembled European DNA sequences. Construct an historical scenario to account for these observations.
20. A magazine article featured parents who filed a “wrongful birth” lawsuit against their doctor for failing to offer prenatal testing for spina bifida, which their daughter was born with in 2003—even though they love the child dearly. They will not say whether they would have ended the pregnancy had they known about the birth defect. If they had ended it, would that have been a eugenic act? Explain your answer.

Forensics Focus

21. In the 1870s, prison inspector and self-described sociologist Richard Dugdale noticed that many inmates at his facility in Ulster County, New York, were related. He began studying them, calling the family the “Jukes,” although he kept records of their real names. Dugdale traced the family back seven generations to a son of Dutch settlers, named Max, who was a pioneer and lived off the land. Margaret, “the mother of criminals,” as Dugdale would write in his 1877 book *The Jukes: A Study in Crime, Pauperism, Disease and Heredity*, married one of Max’s sons, and the couple presumably ultimately gave rise to 540 of the 709 criminals on Dugdale’s watch. Dugdale attributed the Jukes’ less desirable characteristics to heredity.

The Jukes study influenced social scientists to probe other families seemingly riddled with misfits—they were all caucasian, descended from colonial settlers, and poor. Poverty was not seen as an economic problem, but as a reflection of inborn degeneracy, that if left unchecked would cost society greatly.

Dugdale’s book fed the fledgling eugenics movement. In 1911, researchers at the Eugenics Record Office in Cold Spring Harbor described the Jukes’ phenotype

as “feeble-mindedness, indolence, licentiousness, and dishonesty.” The Jukes story and others were used to support compulsory sterilization of those deemed unfit. But the original research on the Jukes family was flawed, and its accuracy never questioned. Less notorious Jukes family members served in respected professions, some even holding public office. The Jukes were vindicated in 2003, when archives at the State University of New York at Albany revealed the original names of the people in Dugdale’s account; most were not even related. The Jukes family curse was more legend than fact.

- a. What would have had to happen to the original jailed Jukes family members or their descendants to be considered eugenic?
- b. How could studies on one family harm others?
- c. Cite an example of an idea based on eugenics today or in the recent past.
- d. If you were a contemporary of Dugdale’s, what type of evidence would you have sought to counter his ideas?



Comparing skulls among modern humans, our modern primate cousins, and fossilized hominins can reveal much about our ancestors and our evolution.

CHAPTER

16

Human Ancestry

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THE HOBBITS

It's odd, in the animal world, to be the only ones of our kind. That wasn't always so, which may be why the theme of a dual humanity persists in science fiction. H. G. Wells' *The Time Machine* looked ahead at two battling breeds of people. In the novel *Darwin's Children*, a virus scrambles the genomes of a group of newborns, starting a new species. Other books describe holdouts from times past—a Neanderthal in modern-day Tajikistan, or a caveman meeting a paleontologist in a Kenyan jungle.

Fossils indicate that from 2 to 6 million years ago, humans and prehumans overlapped, in time if not place. The discovery of preserved bones of several ancient humans on the island of Flores in Indonesia in 2004 suggested a possible real and recent coexistence of two types of people. A female skeleton found 17 feet beneath a cave floor with pieces of others nearby was named *Homo floresiensis*, popularly called the Hobbit. She was about half as tall as a modern human, with a brain about a third of the size. She lived about 18,000 years ago.

The Hobbits exhibited "island dwarfism," an effect of natural selection on small, isolated, island populations of animals. With limited resources, individuals who need less food are more likely to live to reproduce. But who were the Hobbits? At first, some researchers thought that Hobbits were direct descendants of *Homo erectus*, a species that lived before our own. Then, analysis of limb bones of the

skeletons revealed feet and proportions like those of an ape, despite the more human-like skull. This analysis suggests that the Hobbits may have been direct descendants of a primate even older than *Homo erectus*. They evolved in a different direction thanks to the isolation of their island.

16.1 Human Origins

We have sparse evidence of our beginnings—pieces of a puzzle in time, some out of sequence, many missing. Traditionally, paleontologists (scientists who study evidence of ancient life) have consulted the record in the earth's rocks—fossils—to glimpse the ancestors of *Homo sapiens*, our own species. Researchers assign approximate ages to fossils by observing which rock layers fossils are in, and by extrapolating the passage of time from the ratios of certain radioactive chemicals in surrounding rock.

Fossils aren't the only way to peek into species' origins and relationships. Modern organisms also provide intriguing clues to the past in their DNA. Sequences of DNA change over time due to mutation, and on a population level by the forces of nonrandom mating, migration, genetic drift, and, most powerfully, natural selection. The premise of DNA sequence comparisons is that closeness of relationship is reflected in greater similarity of sequence. The logic is that similar sequences are more likely to have arisen from individuals or species sharing ancestors than from the exact same set of spontaneous mutations occurring by chance. By analogy, it is more likely that two young women wearing the same combination of clothes and accessories purchased them at the same store than that each happened to assemble the same collection of items from different sources. On rare occasions, DNA is available from ancient specimens to add to what we know from DNA sequences of modern organisms.

Treelike diagrams are used to depict evolutionary relationships, based on fossil evidence and/or inferred from DNA sequence similarities. Branchpoints on the diagrams represent divergence from shared ancestors. Overall, evolution is shown as a series of branches as species diverged, driven by allele frequencies changing in response to the forces discussed in chapter 15: nonrandom mating, genetic drift, migration, mutation, and natural selection. Evolution is *not* a linear morphing of one type of organism into another—a common misunderstanding (figure 16.1). Humans and chimps diverged from a shared ancestor; humans didn't form directly from chimps. Similarly, two second cousins share great-grandparents, but one cousin did not descend from the other.

In this chapter, we explore human origins, and the genetic and genomic evidence for our evolution.

Hominoids and Hominins

A species includes organisms that can successfully produce healthy offspring only among themselves. *Homo sapiens* (“the wise human”), our species, probably first appeared during the Pleistocene epoch, about 200,000 years ago. Our ancestry reaches farther back, to about 60 million years ago when rodent-like insect eaters flourished. These first primates diverged to give rise to many new species. Their ability to grasp and to perceive depth provided the flexibility and coordination necessary to dominate the treetops.

About 30 to 40 million years ago, a monkeylike animal the size of a cat, *Aegyptopithecus*, lived in the lush tropical forests of Africa. Although the animal probably spent most of its time in the trees, fossilized remains of limb bones indicate it could run on the ground, too. Fossils of different individuals found together indicate that they were social animals. *Aegyptopithecus* had fangs it might have used for defense. The large canine teeth seen only in males suggest that males may have hunted to feed their mates. *Propliopithecus* was a monkeylike contemporary of *Aegyptopithecus*. Both animals are possible ancestors of gibbons, apes, and humans.

From 22 to 32 million years ago, Africa was home to the first **hominoids**, animals ancestral to apes and humans only. One such resident of southwestern and central Europe was called *Dryopithecus*, meaning “oak ape,” because its fossilized bones were found with oak leaves (figure 16.2a). The way the bones fit together suggests that this animal lived in the trees but could swing and walk farther than *Aegyptopithecus*.

More abundant fossils represent the middle-Miocene apes of 11 to 16 million years ago. These apes were about the size of a human 7-year-old and had small brains and pointy snouts. (*Miocene* refers to the geologic time period).

Apelike animals similar to *Dryopithecus* and the middle-Miocene apes flourished in Europe, Asia, and the Middle East during the same period. Because of the large primate population in the forest, selective pressure to venture onto the grasslands in search of food and habitat space must have been intense. Many primate species probably vanished as the protective forests shrank. One type of middle-Miocene ape survived to give rise to humans and African apes. Eventually, animals ancestral to humans only, called **hominins**, arose and eventually thrived. (An older term is *hominid*.)

Hominoid and hominin fossils from 4 to 19 million years ago are scarce, and are often just fragments of tooth and jaw. About 6 million years ago, the hominin lineage split from the apes. There are at least three candidates for this first primate one step closer to humanity from the chimp: *Ardipithecus kadabba* from Ethiopia, *Sahelanthropus tchadensis* from Chad, and *Orrorin tugenensis* from Kenya. They are near the base of the evolutionary tree diagram in figure 16.3, which depicts probable relationships among some of our relatives, past and present. This evolutionary tree is based on fossil evidence and DNA sequence comparisons for the modern species.

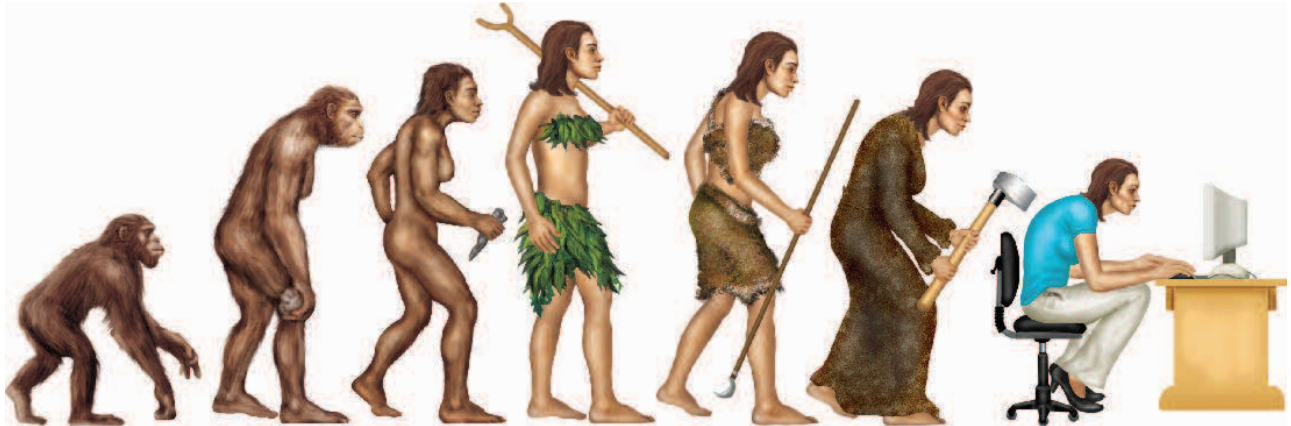


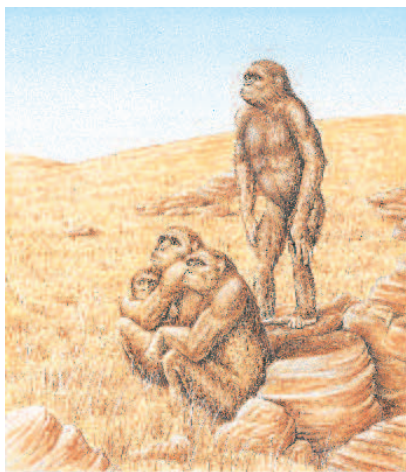
Figure 16.1 Evolution is a branching from shared ancestors. However, the very recent evolution of hominins appears as a single line of descent—one branch—because we are the only modern people. Cartoonists have drawn several variations on the evolutionary theme, often oversimplifying as a straight line. The tree in figure 16.3 is more accurate.

Fossil evidence is more complete for our ancestors who lived 2 to 4 million years ago, who walked upright and conquered vast new habitats on the plains. Researchers apply computer modeling to simulate working versions of ancient body parts to supplement and extend clues from fossils. Several species of a hominin called *Australopithecus* lived at this time, probably following a hunter-gatherer lifestyle. More than one species could coexist because they lived in small, widely separated groups that probably never came into contact. The australopithecines were gradually replaced with members of our own genus, *Homo*. The following sections introduce a few of these ancestors known from rare fossil remains and what our imaginations can fill in.

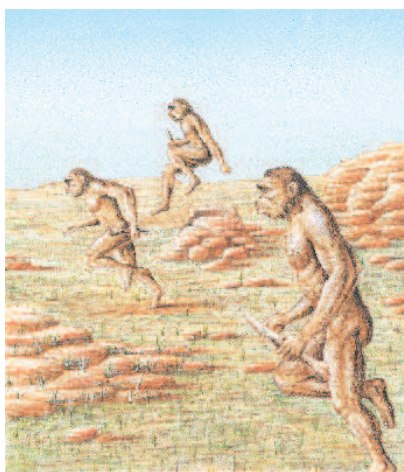
Australopithecus

Peoples of the Past—The Dikika Infant, 3.3 Million Years Ago

The 3-year-old had probably wandered away from her family when she had been swept up in the sudden flood 3.3 million years ago in Ethiopia. She liked to sit up in the trees ringing the large lake to watch the antelopes and giraffes, elephants and wildebeests. She wandered the grasslands to the small rivulets that merged into the Awash river delta, avoiding crocodiles while she waded among the reeds, picking up snails and trying to catch fish. One day, perhaps after seasonal rains, the



a. *Dryopithecus*



b. *Australopithecus*



c. *Homo erectus*

Figure 16.2 Human forerunners. (a) The “oak ape” *Dryopithecus*, who lived from 22 to 32 million years ago, was more dextrous than his predecessors. (b) Several species of *Australopithecus* lived from 2 to more than 4 million years ago, and walked upright. (c) *Homo erectus* made tools out of bone and stone, used fire, and dwelled communally in caves from 35,000 to 1.6 million years ago.

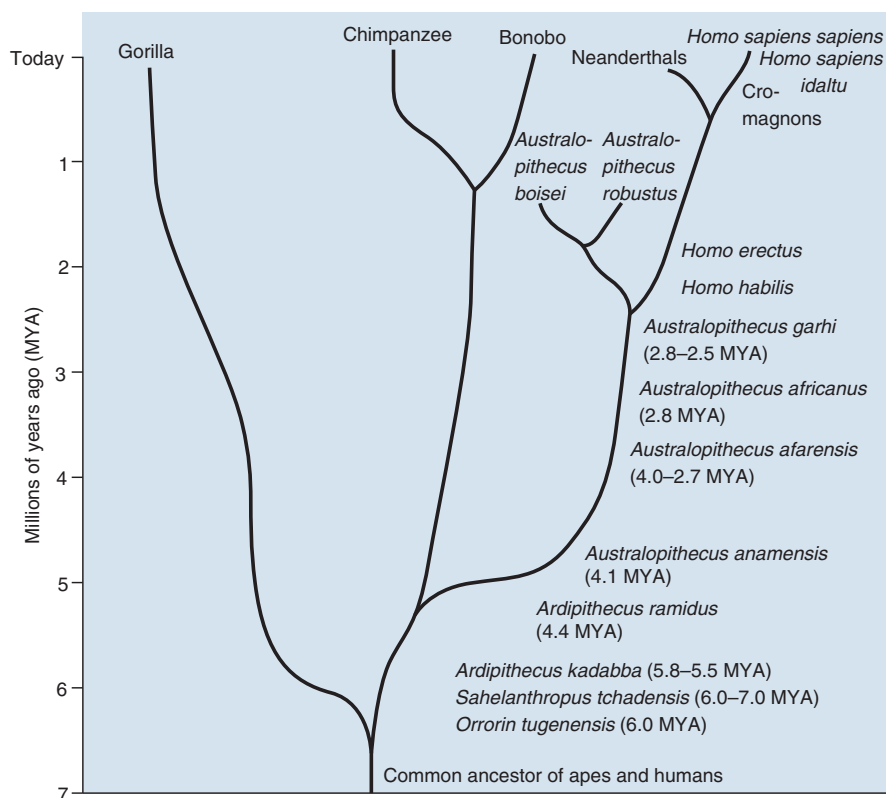


Figure 16.3 Evolutionary tree. An evolutionary tree diagram indicates the relationships among primates, past and present. (Not all known hominin species are indicated.)

girl was overcome by rushing waters and was quickly buried in thick sediments. Much of her skeleton remained intact.

Paleontologists discovered the top of her skull protruding from sandstone. After 5 years of meticulously removing the encasing stone with dental instruments, clues to who the Dikika infant was began to emerge. The tiny, delicate bones of the child, representing *Australopithecus afarensis*, reveal a mix of characteristics of human and chimp. Her legs were shorter but her arms were longer than ours. Her upper half looked like that of an ape, with a large, protruding lower face and neck muscles to support it. She was hairy, with arms long enough to have easily grabbed a branch, her hands and fingers curved enough to grasp and hold on. Yet her bottom half was much more like that of a human. The great toes aligned with the other toes, the hip joints and leg bones able to support walking, although hunched because the knee joints could not lock. She had a voice box, and while some of her teeth resembled those of apes, some were more humanlike. Her cranium was slightly larger than that of a chimp. She looked like an ape but walked like a human.

Australopithecines had flat skull bases, as do all modern primates except humans. They stood about 4 to 5 feet tall. The angle of preserved pelvic bones, plus the discovery of *Australopithecus* fossils with those of grazing animals, indicate that this ape-human had left the forest.

The oldest species of australopithecine known, *Australopithecus anamensis*, lived about 4.1 million years ago. A partial skeleton, named Lucy, represents an individual who lived about 3.6 million years ago in the same area as the Dikika infant—the Afar river basin of Ethiopia (figure 16.4). This is the only place where fossil evidence of our ancestors spans 6 million years. Lucy died, with arthritis, at about age 20.

Other fossils offer additional clues to australopithecine life. Two parallel paths of humanlike footprints, preserved in volcanic ash in the Laetoli area of Tanzania, are contemporary with Lucy. A family may have left the prints, which are from a large and small individual walking close together, with a third following in the steps of the larger animal in front. Researchers at Arizona State University used computer simulation with measurements from modern monkeys and chimps as well as from a fossilized ancestor of Lucy to reveal that the australopithecine jaw was strong enough to crack hard nuts—an adaptation to a changing environment.

Toward the end of the australopithecine reign, *Australopithecus garhi* may have coexisted with the earliest members of *Homo*. Its fossils from the Afar region date from about 2.5 million years ago. Remains of an antelope found near the australopithecine fossils suggest butchering. The ends of the long bones had been cleanly cut with tools, the marrow removed, meat stripped, and the tongue cleanly sliced off.



a.



b.

Figure 16.4 Australopithecus. Lucy's skeleton offers many clues to what this hominin was like (a). About 3.6 million years ago, she walked upright in the grasses along a lake in the Afar region of Ethiopia, about six miles from where the Dikika infant would live 300,000 years later. She skimmed the shores for crabs, turtles, and crocodile eggs to eat. (b) An artist's interpretation of what Lucy may have looked like.

A. garhi stood about 4.5 feet tall, and like the Dikika infant and Lucy, the long legs were like those of a human, but the long arms were more like those of an ape. The small cranium and large teeth hinted at apelike ancestors.

Homo

Peoples of the Past—Idaltu Man, 156,000 Years Ago

In 1997, paleoanthropologist Tim White was driving by the village of Herto, along a bend of the Awash River. Seasonal rains had driven the nomadic people and their livestock away, and had cleared the ground in places. In one such bald spot, White spotted a hippopotamus skull sticking up. Near it were tools

*made of obsidian, a glasslike rock. A few days later he sent two students to explore further, and they found a humanlike skull lying on its side. Soon, two other skulls were discovered. One was from another adult, and the other was a child's skull shattered and scattered into more than 200 pieces, including baby teeth. The researchers named the hominin *Homo sapiens idaltu*, which in the local language means "elder."*

The most intact skull was slightly longer, and the brain slightly larger, than those of modern humans. Fine, parallel lines had been etched along the base of the skull. The dome of the skull had not been damaged, as it would have been had cannibalism been practiced. The skull was very smooth, and there were no other bones nearby. Might the skulls have been gently

separated from the bodies, saved, and touched, as modern cultures do to honor the dead?

Other fossils filled in the story. Evidence of catfish and hippos indicate that the Awash River had flooded, forming a freshwater lake. The hippo and buffalo bones bore marks made with tools that had probably sliced off meat. Some bones were broken in ways that suggested that the people ate the marrow. The tools were of a sophisticated design compared to the flaked tools from a million years ago. Overall, the scene evoked an image of ancestors who not only understood the concept of death, but who practiced mortuary rituals.

Our knowledge of how *Homo* replaced *Australopithecus* is sparse. Some australopithecines were “dead ends” that died off. Clues suggest that by 2.3 million years ago, *Australopithecus* coexisted with *Homo habilis*—a more humanlike cave dweller who cared intensively for its young. *Habilis* means handy, and this primate was the first to use tools for tasks more challenging than stripping meat from bones. *H. habilis* may have descended from hominins who ate a greater variety of foods than other ape-humans, allowing them to live in a wider range of habitats.

H. habilis coexisted with and was followed by *Homo erectus* during the Paleolithic Age (table 16.1). One famed *H. erectus* fossil, named “Daka” for the place where it was found in the Afar region, is from an individual who lived about a million years ago. It had a shallow forehead, massive brow ridges, a brain about a third smaller than ours, and strong, thick legs. Daka lived on a grassland, with elephants, wildebeests, hippos, antelopes, many types of pigs, and giant hyenas. Figure 16.5 depicts what he might have looked like.

H. erectus left fossil evidence of cooperation, social organization, tools, and use of fire. Fossilized teeth and jaws suggest that they ate meat. The distribution of fossils indicates that they lived in families of male-female pairs (most primates have harems). The male hunted, and the female nurtured the young. They were the first to have an angled skull base that permitted them to produce a greater range of sounds, making speech possible. *H. erectus* fossils have been found in China, Java, Africa, Europe, and southeast Asia, indicating that these animals could migrate farther than earlier primates.

The fossils of *H. sapiens idaltu* from Ethiopia reveal that by 156,000 years ago, our ancestors did not look very different from ourselves (figure 16.6). *H. sapiens idaltu* probably resembled an Australian aborigine, with a large and powerful build and dark skin.

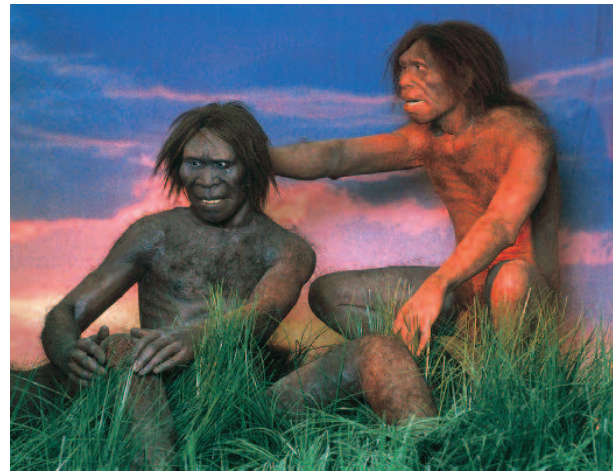


Figure 16.5 *Homo erectus*. This artist's rendition is based on clues from many fossils.

By 70,000 years ago, humans, still mostly confined to Africa, used more intricately carved tools made of bones, and red rock that bore highly symmetrical hatchmarks, which may indicate early counting. Groups of hominins may have been very isolated on the vast continent. The first *Homo* may have left Africa around 100,000 years ago, as a founder group of about a thousand individuals, most of them male, who died out by 70,000 years ago. Because of the isolation, it's possible that even as *H. sapiens idaltu* and perhaps others yet to be discovered were far along the road to modern humanity, pockets of *H. erectus* may have persisted, perhaps until as recently as 35,000 years ago.

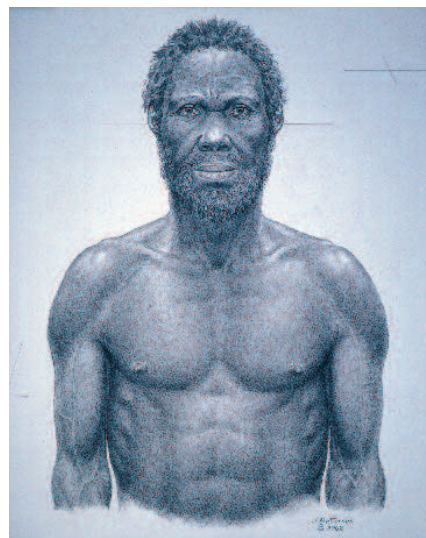


Figure 16.6 *Homo sapiens idaltu*. Discovery of three skulls made possible this artist's depiction of what this early member of our species might have looked like—not very much different from us.

Table 16.1 Cultural Ages

Age	Time (years ago)	Defining Skills
Paleolithic	750,000 to 15,000	Earliest chipped tools
Mesolithic	15,000 to 10,000	Cutting tools, bows and arrows
Neolithic	10,000 to present	Complex tools, agriculture

A find of nine Neanderthal skeletons in a cave in northern Spain called El Sidron has contributed much to our knowledge of these people. In addition to the skeletal characteristics described in **figure 16.7**, the remains had sharply broken bones, indicating cannibalism to eat the marrow. The Neanderthals (*Homo neanderthalensis*) were also contemporaries of *H. erectus* and members of genus *Homo*. Our knowledge of the Neanderthals comes from fossil evidence as well as analysis of DNA sequences from that evidence, discussed in the next section. The Neanderthals split from the lineage that led to us about 516,000 years ago. A founding group of about 3,000 Neanderthals left Africa, traveling slowly toward Europe where, by about 150,000 years ago, they were widespread. Neanderthals and modern people may have coexisted in what is now Israel about 90,000 years ago. By 70,000 years ago, the Neanderthals had spread to eastern Asia, even reaching Siberia. They lived all over Europe, probably never numbering more than about 15,000 individuals.

Fossil evidence indicates that from 30,000 to 40,000 years ago, the Neanderthals coexisted with the lighter-weight, finer-boned Cro-Magnons. The newcomers had high foreheads and

well-developed frontal brain regions, and signs of culture that we do not see for Neanderthals. The first Cro-Magnon fossils were found in a French cave. Five adults and a baby were arranged in what appeared to be a communal grave. Nearby were pierced seashells that may have been used as jewelry. Intricate art decorated the cave walls. In contrast, the few Neanderthal graves show no evidence of ritual, just quick burial.

The most recent Neanderthals that we know of lived from 28,000 to 24,000 years ago in at least eight settlements on Gibraltar. This area is the southernmost part of Europe, and remained warm when the rest of the continent underwent ice ages. Caves there had rocky ledges that may have served as hearths, as well as high ceilings that let in light. Tools left there suggest that Neanderthals may have occupied these cozy domains, possibly on and off, for 100,000 years or longer. They shared Gibraltar with the same types of animals seen on the warm, dry Serengeti plain of Africa, and also with bears, wolves, deer, cattle, ducks, tortoises, and pine and olive trees. Food was abundant, the climate warm, and the caves protective.

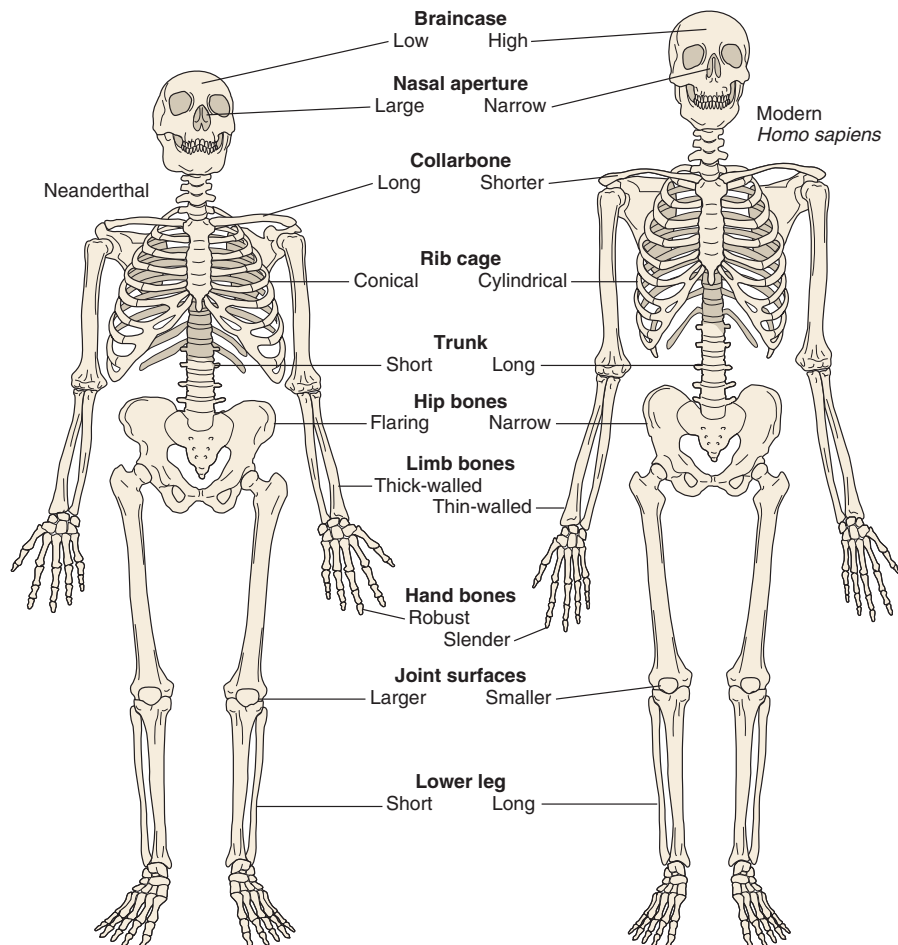


Figure 16.7 Neanderthals. Neanderthals probably looked a lot like us, but there were many subtle skeletal differences. We share ancestors with Neanderthals, but they are not our direct ancestors. They are a side branch on our family tree.

The decline of the Neanderthals may have been due to competition with the lighter and presumably smarter archaic humans, and because their large bodies simply did not enable them to migrate fast enough to escape the cold. The fact that Neanderthals lived until as recently as 24,000 years ago, coupled with evidence that our direct ancestors lived in Europe by 32,000 years ago, suggests a timing that might have allowed breeding between the two groups. But DNA sequence evidence suggests that even if interbreeding happened occasionally, it was too rare to have left a mark in the modern human genome.

The Neanderthals take their name from Neander Valley, Germany, where quarry workers blasting in a limestone cave on a summer day in 1856 discovered the first preserved bones. A Neanderthal discovered in France fifty years later, the “Old Man” of La Chapelle-aux-Saints, led to their common depiction as primitive and slow-witted, stooped perhaps due to arthritis. Since then, reconstructions of other Neanderthal skeletons indicate major differences from us.

Compared to a modern human, a Neanderthal had a wider pelvis, shoulders, and ribcage, and shorter forearms and shins; prominent brow ridges, a forward-pointing face, and a sloping forehead. The characteristic heavy brow bones might have resulted from genetic drift on populations isolated in cave systems. A fossilized, deformed skeleton buried with flowers in Shanidar Cave, Iraq, reveals that the Neanderthals may have been spiritual hunter-gatherers. We return to the Neanderthals later in the chapter.

Modern Humans

Peoples of the Past: Ötzi

In 1991, hikers in the Ötztal Alps of northern Italy discovered an ancient man frozen in the ice (**figure 16.8**). Named Ötzi, the Ice Man was on a mountain more than 10,000 feet high 5,200 years ago when he perished. He wore furry leggings, leather suspenders, a loincloth, fanny pack, bearskin cap and cape, and sandal-like snow shoes. Berries found with him place the season as late summer or early fall. His last meal was ibex and venison.

Ötzi died following a fight. He had a knife in one hand, cuts and bruises, and an arrowhead embedded in his left shoulder that nicked a vital artery. The wound bore blood from two other individuals, and his cape had the blood of a third person. Mosses found on his body may have been wound dressings. He likely bled to death, falling into a ditch, where he was covered by snow. After this safe burial, which preserved his body intact, a glacier sealed the natural tomb. DNA profiling suggests that he belonged to the same gene pool as modern people living in the area, which is near the Italian-Austrian border.

Cave art from about 14,000 years ago indicates that by that time, our ancestors had developed fine-hand coordination and could use symbols—milestones in cultural evolution. By 10,000 years ago, people had expanded from the Middle East across Europe, bringing agricultural practices.

Another way that anthropologists try to glimpse what humans were like a few thousand years ago is by studying vanishing indigenous peoples, such as the San (bushmen) and Pygmies of Africa, the Etas of Japan, the Hill People of New



Figure 16.8 A 5,200-year-old man. (a) Hikers discovered Ötzi, the Ice Man, in the Austrian/Italian Alps in 1991. He lived 5,200 years ago. (b) Ötzi wore well-made clothing, including a hat; used intricate arrows that demonstrate familiarity with ballistics and engineering; and carried mushrooms with antibiotic properties. He had tattoos, indentations in his ears that suggest he wore earrings, and evidence of a haircut. This depiction is derived from the evidence found on and near Ötzi's preserved body.

Guinea, and a Brazilian tribe, the Arawete, who number only 130 individuals. Studying DNA sequences in these populations provides information on their origins because the people have stayed in the same geographical region and not mixed their gene pools with others. However, researchers wishing to sample DNA from these groups have run into problems when science clashes with culture. In some indigenous societies, ancestry is extremely important, and provides the basis for identity and rights. Some people fear that genetic research might be used to dispute ancestry claims. *Bioethics: Choices for the Future* takes a closer look at indigenous peoples.

Figure 16.9 is a timeline representing some of the human ancestors discussed in this chapter. The rest of the chapter examines how DNA sequences can shed light on the past, and how humanity has populated the Earth.

Key Concepts

1. Monkeylike *Aegyptopithecus* lived about 30 to 40 million years ago and was ancestral to gibbons, apes, and humans. The first hominoid, *Dryopithecus*, lived 22 to 32 million years ago and may have walked onto grasslands.
2. Hominins appeared about 19 million years ago.
3. About 4 million years ago, bipedalism opened up new habitats for *Australopithecus*. *A. garhi* may have coexisted with the first *Homo*.
4. By 2 million years ago, *Australopithecus* coexisted with *Homo habilis*. Later, *H. habilis* coexisted with *H. erectus*, who used tools in more complex societies. *H. erectus* then coexisted with *H. sapiens*. *H. sapiens idaltu* lived 156,000 years ago.
5. The Neanderthals were a side branch from modern humans who disappeared about 30,000 years ago.
6. A preserved man from 5,200 years ago is genetically like us.

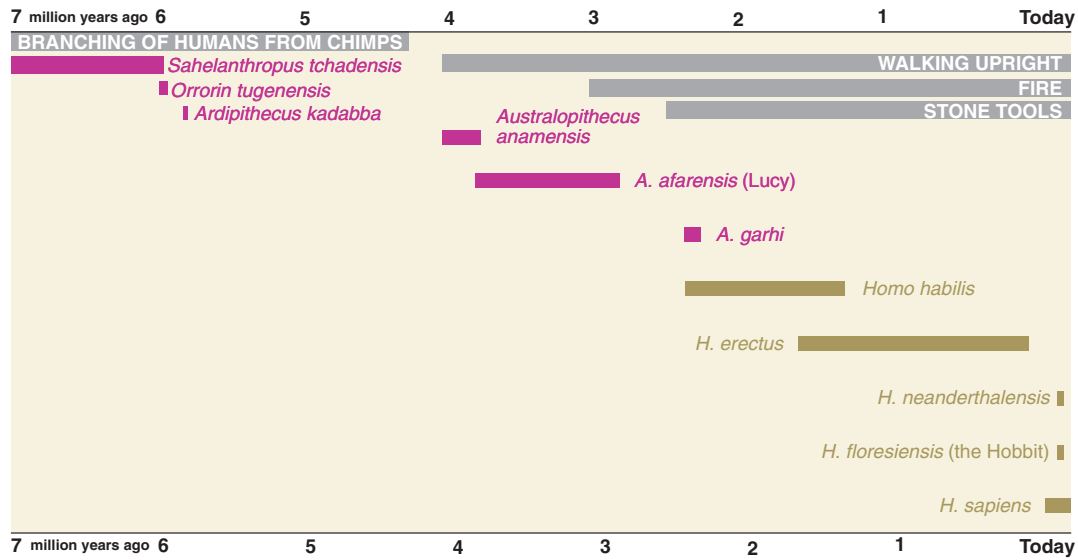


Figure 16.9 Our most recent ancestors.

Bioethics: Choices for the Future

Indigenous Peoples Provide Clues to the Past

Not everyone on the planet uses cell phones, computers, and electricity, or gets food at a supermarket. Living much closer to the land, sometimes not much different from our hunter-gatherer ancestors, are 380 million or so members of indigenous peoples, who account for less than 6 percent of the world's population.

An indigenous group, by definition, can trace its ancestry back farther in a particular geographic area than any other group, and has retained its uniqueness in cultural practices, social organization, and/or language. The group has remained physically isolated, or culturally isolated, while surrounded by colonists. Because of this isolation, indigenous groups have also kept separate their gene pools, which may be quite different from those of their neighbors.

About 5,000 indigenous peoples are known, but others may exist. They have been identified in more than 72 nations, and range from just a few dozen individuals to making up sizeable portions of a country's population. Some live in distinct tribes, such as the Maori of New Zealand. In some nations indigenous peoples have been marginalized, and have poor health and live in poverty. The accompanying table lists some indigenous peoples.

Indigenous peoples are threatened, with either extinction or assimilation, as the rest of the world continues to expand. Efforts to understand their way of life often go awry. This was the case for the Huaorani Indians of Ecuador and Peru, who live in the Amazon rainforest (see photo, top). They were unknown until an oil company showed up to investigate drilling for oil in the late 1940s, and left when threatened by the Huaorani's spears and poisoned darts. A few years later, missionaries were killed. "Huaorani" means "human being," and the people refer to all others as "cowode," or "nonhuman." They are

Indigenous Peoples	
People	Home
Adivasis	India
Basque	France/Spain
Chamorros	Guam and northern Marianas
Huli	Papua New Guinea
Inuit	Canada and Greenland
Kalahari	Botswana
Komi	Western Urals
Kurds	Iran
Malagasy	Madagascar
Maori	New Zealand
Sami	Northern Scandinavia
Yanomami	Brazil and Venezuela

hunter-gatherers who lead a semi-nomadic lifestyle, moving about every 10 years. We know little about their genetics, but we do know that they suffer from infections brought in by the missionaries, and skin disorders, hair loss, and sore throats thanks to pollution from the oil industry.

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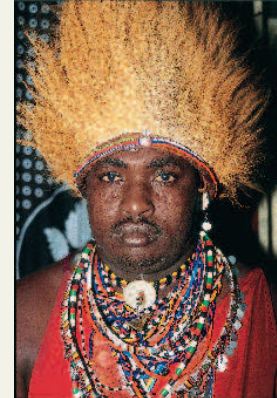


A Huaorani Indian holding a spear.

Geneticists have attempted to describe the genetic diversity among nine indigenous groups in Africa, but so far have not been able to distinguish them. It may take more diverse SNP markers to differentiate these populations whose roots may go back much farther than those of other peoples. The man in the photo (right) is a chief of a Maasai tribe, who live in Kenya and Tanzania and are one of the nine groups being studied. The Maasai have low body weight and blood pressure, and excellent lipid profiles, despite a very fatty diet. Their healthy hearts and blood vessels are likely due more to their very active nomadic lifestyle than to inheriting protective gene variants, researchers think.

Studying indigenous peoples has been controversial. The most celebrated case was that of the Yanomami, who live near streams in southern Venezuela and northern Brazil. In 1968, a team of anthropologists led by geneticist James Neel gave the Yanomami a measles vaccine that they

knew to be dangerous to people who had had no contact with the outside world. Hundreds of Yanomami died, as Neel prevented anyone from helping them—he supposedly wanted to observe the effects of introducing an infectious disease to an isolated group. He died in 2000, the truth never fully revealed. But the controversy has not died. Blood was sampled from the Yanomami for comparison to other modern populations to assess genetic diversity, but the people were not informed that the samples might be used in research after their deaths. The Yanomami religion forbids keeping tissue after death. The controversy continues as the people attempt to take back their blood samples from research labs.



This Maasai chief wears the traditional headware and accessories.

The controversy continues as the people attempt to take back their blood samples from research labs.

Questions for Discussion

1. Discuss why geneticists would want to study the DNA of indigenous peoples.
2. Under what conditions do you think it is ethical for geneticists to study the DNA of indigenous peoples?
3. Journalist Patrick Tierney wrote a book, *Darkness in El Dorado*, that chronicles James Neel's work with the Yanomami. Read reviews of the book and discuss what the research team did, and whether or not you think their actions were justified.

16.2 Molecular Evolution

Fossils paint an incomplete picture of the past because they are scarce and only certain parts of certain organisms were preserved. Additional information comes from the informational molecules of life, DNA and amino acid sequences, which change in sequence over time as mutations occur. The more alike a gene or protein sequence is in two species, the more closely related the two are presumed to be—that is, the more recently they shared an ancestor. The assumption is that it is highly unlikely that two unrelated species would evolve precisely the same sequence of DNA nucleotides by chance.

Comparing genome, DNA or protein sequences, and chromosome banding patterns constitute the field of **molecular evolution**. Knowing the mutation rates for specific genes provides a way to measure the passage of time—a sequence-based molecular clock, of sorts. Sequence information was used to build the evolutionary trees in **figure 16.10**.

Comparing Genes and Genomes

We can assess similarities in DNA sequences between two species for a piece of DNA, a single gene, a chromosome segment, a chromosome, mitochondrial DNA, or an entire genome.

For some genes, similarities among species can be startling. People with Waardenburg syndrome (MIM 148820), for example, have a white forelock of hair; wide-spaced, light-colored eyes; and hearing impairment (**figure 16.11**). The mutant gene is very similar in sequence in cats, horses, mice, and minks, who have light coats and eyes and are deaf. In these species, the phenotype stems from abnormal movements of pigment cells in the embryo's outermost layer.

In general, DNA sequences that encode protein are often very similar among closely related species. The related species presumably inherited the gene from a shared ancestor, and a change in that gene would not persist in a population unless it provided a selective advantage. At the same time, natural selection weeded out proteins that did not promote survival to reproduce.

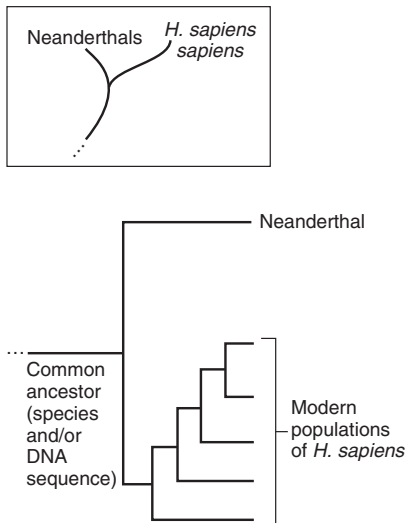


Figure 16.10 Neanderthals split from the lineage that led to *Homo sapiens*, as this molecular evolutionary tree diagram indicates. The inset shows the corresponding information on the stylized version of a tree in figure 16.3.

Similar DNA or amino acid sequences in different species are said to be “highly conserved.” Sequences that are similar in closely related species but that do not encode protein often control transcription or translation, and so are also vital and therefore subject to natural selection. In contrast, some genome regions that vary widely among species do not affect the phenotype, and are therefore not subject to natural selection. Within a protein-encoding gene, the exons tend to be highly conserved, but the introns, which are removed from the corresponding RNA, are not.

⊕ Solving a Problem

Comparing Chimps and Humans

In many ways we have more in common with chimpanzees than with any other animal. Just how similar we are at the genome level, however, depends upon how we assess

similarity—by DNA sequence, numbers of copies of sequences, or sequences missing from the human genome. The commonly repeated estimate of 98.7 percent sequence similarity at the genome level between human and chimp originated from studies conducted in the 1970s using a technique called DNA hybridization. DNA from two species is unwound, cut, and mixed. Complementary pieces bind, and some hybrid molecules form, with one strand of the double helix from one species, the other from the other species. The premise is that the higher the temperature required to separate hybrid double helices, the more of the sequence is shared, because more complementary base pairs attract (figure 16.12). These studies used DNA segments present in single copies in the genome, which indicates that they likely encode protein. Protein comparisons support the 98.7 percent sequence identity, too.

Another way that the genomes of human and chimp differ is in the number of copies of certain DNA sequences. Genome similarity can be estimated by tracking “indels,” which stands for “insertions and deletions.” Compared to the chimp genome, the human genome has additional copies of certain sequences (“insertions”) and lacks certain sequences (“deletions”). If small insertions and deletions that distinguish the human and chimp versions of the same gene are considered, then our degree of genome similarity is only about 96.6 percent. The degree of similarity may even be as low as 94 percent if sequences not in the human genome are considered. That is, what *isn't* present defines us as well as what *is* present.

A simple calculation on a short, hypothetical DNA sequence demonstrates the effect of small differences. Consider an ancestral sequence of 15 bases:

G A T A C G A G C T C T A A C

“Ancestral” means that the most recent common ancestor of humans and chimps had this sequence. If a single-base substitution occurred after the divergence from the shared ancestor, then the correspondence would be less than 100 percent:

Chimp G A T A C G A G C T C T A A C
Human G A T A C G A G C T A T A A C



a.



b.



c.

Figure 16.11 The same mutation can cause similar effects in different species. A mutation in mice (a), cats (b), humans (c), and other types of mammals causes light eye color, hearing or other neurological impairment, and a fair forelock.

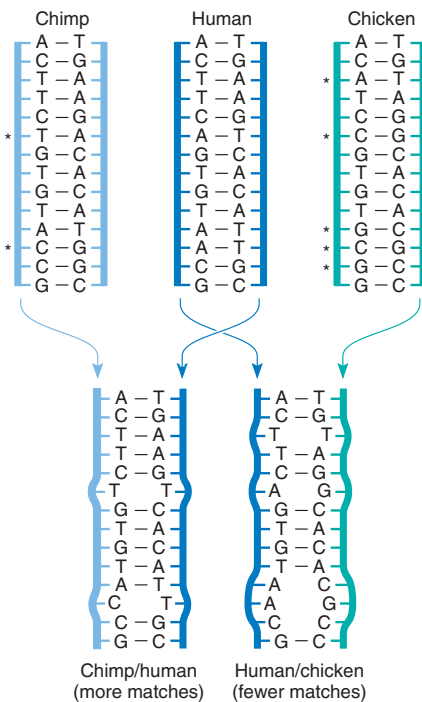


Figure 16.12 The rate of DNA hybridization reflects the degree of evolutionary relatedness. This highly schematic diagram shows why DNA from a human hybridizes more rapidly with chimpanzee DNA than with chicken DNA. Each * refers to a site where chimp or chicken DNA differs from human DNA.

After the C-to-A point mutation occurs, humans and chimps share 14 of these 15 bases, for a correspondence of 93.3 percent, rather than 100 percent identity. But what happens when three bases at a time (so as not to offset the reading frame) are added or deleted in one of the evolutionary lines? Imagine that three bases inserted into the human lineage as follows:

```
Chimp  GATACGAGCTCTAAC
Human  GATGCAACGAGCTCTAAC
```

The correspondence is now 15 out of 18 bases that are the same, or 83.3 percent.

If the human lineage lost three bases, the correspondence would also diminish:

```
Chimp  GATACGAGCTCTAAC
Human  GATAGCTCTAAC
```

The sequence now shares 12 out of 15 bases, or 80 percent.

The similarity between the human and chimp genome decreases even more if noncoding regions, such as introns and repeats, are considered. However, differences in indels and repeats do not explain how we differ from chimps on a whole organism level. To assess phenotypic distinctions, it is more helpful to look at individual traits, some of which are determined by single genes. Individual genes can

actually have great effects on appearance, physiology, and development.

Genes That Help to Define Us

Uniquely human traits include spoken language, abstract reasoning ability, highly opposable thumbs, and larger frontal lobes of the brain (**Reading 16.1**). One stark difference between chimp and human that could stem from a single gene is hairiness. Chimpanzees and gorillas express a keratin gene whose counterpart in humans is not expressed. When our ancestors left the forests, natural selection might have favored loss of body hair to provide more efficient cooling or as a way to shed skin parasites such as lice. Speech may also be due to a single gene difference between humans and chimps. A family in London whose members have unintelligible speech led to the discovery of a single gene (*FOXP2*, MIM 605317) that controls speaking ability—and is present, but different, in chimps.

Another single gene that accounts for great differences among primates controls the switch from embryonic to fetal hemoglobin (see figure 11.2). More primitive primates lack or have very little fetal hemoglobin. In more recently evolved and more complex primates, fetal hemoglobin correlates to lengthened fetal period, which extended the time for brain growth. With larger brains came greater skills. Single genes can also explain the longer childhood and adolescence in humans compared to chimpanzees.

Single genes that distinguish humans from chimps appear to be few, but they tend to be implicated in Mendelian disorders. Perhaps this reflects the fact that the genes that distinguish us have recently taken on their new functions, and the genome has not yet had time for protective redundancies to have evolved.

The major reason why humans and chimps are genetically so similar, but look and behave so differently, is differences in gene expression, not genome sequence. For example, a study contrasted gene expression in the liver and brain in the two species. The differences in the brain were far greater than in the liver. It makes sense that our livers are more alike than our brains (or at least we'd like to think so!).

Comparisons of the human genome sequence to those of other species are interesting, too (see figure 22.8). Our close relationship to the other vertebrates is revealed by comparing the human genome sequence to that of the pufferfish *Tetraodon nigroviridis*. Its genome is like ours, minus many of the repeats and introns. It is odd to think that the protein-encoding portion of our genome is nearly the same as that of a fish.

Considering Genomes

Overall, the human genome has a more complex organization of the same basic parts as the fruit fly and roundworm genomes. For example, the human genome harbors thirty copies of the gene that encodes fibroblast growth factor, compared to two copies in the fly and worm genomes. This growth factor is important for the development of highly complex organs.

Reading 16.1

What Makes Us Human?

Comparison of the chimpanzee and human genomes has revealed “human accelerated regions.” These are highly conserved sequences (genes present in humans and apes) that show signs of positive selection in humans, such as an amino acid change seen in all human groups but not in the chimp or orangutan versions of the same gene. These genes may represent characteristics that in some way distinguish us from our closest primate relatives. Signs of positive selection in the human genome flesh out views of our ancestry from paleontology.

Tool Use

A paleontologist enjoys a different view of the origin of humanity than that of a geneticist. University of California, Berkeley, paleontologist Tim White heads a team that, for months at a time, calls the Afar region of Ethiopia home. Here, scattered and at different levels, lie remains of our ancestors stretching back some 6 million years, teasing at the time when our forebears split from an ancestor shared with the chimpanzee.

White led the teams that discovered *Ardipithecus*, *Australopithecus garhi*, “Daka,” and *H. sapiens idaltu*. He sums up what distinguishes our species in one word: culture. His imagination takes him back in time:

It all started about 2.5 million years ago, when guys started banging rocks together. That’s what allowed the niche to expand in the beginning, the start of culture. Tool making, utilizing stone, probably began in *Australopithecus*, such as Lucy. They were very adaptable and very widespread, all over Africa. These bipeds were small-brained, and they weren’t busy becoming human, but being australopithecines. At some point, a population of that highly intelligent, bipedal generalist organism that was *Australopithecus* began to exhibit behaviors that we see in the chimp. Chimps hunt monkeys. But chimps lack tools. At some point, an early hominid didn’t lack those tools anymore, and that particular sect formed the beginning of the lineage that would ultimately diverge from other australopithecines that kept on being australopithecines, and that lineage would go on to become early *Homo*. There may have been different varieties, but eventually there was *Homo erectus*.

Walking

Diseases of modern humanity can reveal traits of evolutionary import. Consider Joubert syndrome (MIM 608629). In this disorder, nerve cell fibers cannot cross from their origin on one side of the brain to the other, so a person cannot move just one arm or leg. In response to a command to move one limb, both move. The part of the brain that controls posture, balance, and coordination is compromised. The gene that causes Joubert syndrome, called *AHI1*, is identical in all modern human groups examined, but has different alleles in chimps, gorillas, and orangutans. Perhaps in the lineage leading to humans, the gene came to control walking by making it possible to place one foot in front of the other.

Running

Homo erectus distinguished itself in another key way: It could run for long distances, thanks to specific anatomical adaptations. The nuchal ligament that connects the skull to the neck became more highly developed in *H. erectus*, enabling the head to stay in place with the force of running. The leg muscles were also more highly developed than those of chimps or australopithecines, acting as springs. *H. erectus* originated a large buttocks, whose muscles contract during running. All three of these structures are not merely the result of being able to walk, but enabled early *Homo*, and us, to run. This skill would have helped our ancestors to escape predators, find food, and locate new homes.

A Big Brain

The difference between a big-brained human and a small-brained chimp may be a few single genes. About 2.4 million years ago, a gene called *MYH16* underwent a nonsense mutation, which prevented production of a type of muscle protein called a myosin. The mutation is seen in all modern human populations, but not in other primates. Without this particular type of myosin, jaw development is not as great. With a diminished jaw, the bony plates of the skull could expand, allowing greater brain growth (**figure 1**). Researchers nicknamed the mutation RFT, for “room for thought.” Fossil evidence indicates that the switch from “big jaw, small brain” to “small jaw, big brain” happened when *Homo* gradually replaced *Australopithecus*, about 2 million years ago. The genetic analysis may be new, but the idea wasn’t. Charles Darwin wrote in 1871 that different-sized jaw muscles were at the root of the distinction between apes and humans.

Cognition

At the genetic level, humans and chimps may differ more in the numbers of copies of particular genes than in the nature of the genes. Researchers identified 134 genes with an increased copy number in the human genome compared to the genomes of the great apes. Many of these genes are involved in brain structure or function. Some of the genes promote the signal transduction that underlies long-term memory; others, when mutant, cause mental retardation or impair language skills. Single genes implicated in fueling human brain growth control the migration of nerve cells in the front of the human fetal brain.

Sense of Smell

Natural selection retains useful traits and weeds out harmful or useless ones. Both of these forces have fine-tuned our “chemical” senses of taste and smell, actually dulling them as our reliance on them for survival waned.

(Continued)

(Continued)

The sense of smell derives from a one-inch-square patch of tissue high in the nose that consists of 12 million cells that bear odorant receptor (OR) proteins. (In contrast, a bloodhound has 4 billion such cells!) Molecules given off by something smelly bind to combinations of these receptors, which then signal the brain in a way that creates the perception of an associated odor.

Our odorant receptor genes number 906, comprise about 1 percent of the genome, and occur in clusters. About 60 percent of them are pseudogenes—their sequences are similar to those of functional “smell” genes, but are riddled with mutations that prevent translation of complete proteins. Perhaps they are remnants of a

distant past when we depended more upon our chemical senses for survival. Natural selection may have, over time, eliminated OR genes no longer essential to survival. Yet other genetic evidence indicates that natural selection also has acted positively to retain the OR genes that continue to function. While the pseudogenes harbor many diverse SNPs (sites where more than one base is common), the functional OR genes are remarkably alike in sequence. In addition, the nucleotide differences that persist among the retained genes actually alter the encoded amino acid, suggesting that natural selection favored these particular sequences.

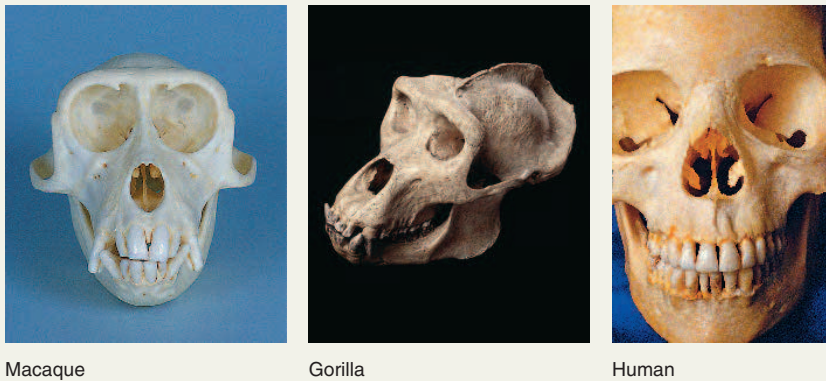


Figure 1 A mutation in a single gene may have made the expansion of the human brain possible.

Genome studies indicate that over deep evolutionary time, genes and gene pieces provided vertebrates, including humans, with certain defining characteristics:

- complex neural networks
- blood clotting pathways
- acquired immunity
- refined apoptosis
- greater control of transcription
- complex development
- more intricate signaling within and among cells

Did the Human Genome Duplicate?

Comparing the human genome to itself provides clues to evolution, too. The many duplicated genes and chromosome segments in the human genome suggest that it doubled, at least once, since diverging from that of a vertebrate ancestor about 500 million years ago. Either the human genome doubled twice, followed by loss of some genes, or, more likely, one doubling was followed by additional duplication of certain DNA sequences.

The extensive duplication within the human genome distinguishes us from other primates. Some of the doublings

are vast. Half of chromosome 20 repeats, rearranged, on chromosome 18. Much of chromosome 2's short arm reappears as almost three-quarters of chromosome 14, and a block on its long arm is echoed on chromosome 12. The gene-packed yet tiny chromosome 22 includes eight huge duplications and several gene families. However, the many repeated DNA sequences in the human genome should not be considered “junk.” Duplications provide raw material and flexibility for future evolution. A copy of a DNA sequence can mutate, allowing a cell to “try out” a new function while the old one carries on. More often, the twin mutates into a silenced pseudogene, leaving a ghost of the gene behind as a similar but untranslated DNA sequence.

A duplication can be located near the original DNA sequence it was copied from, or away from it. A sequence repeated right next to itself is called a tandem duplication, and it usually results from mispairing during DNA replication. A copy of a gene on a different chromosome may arise when messenger RNA is copied (reverse transcribed) into DNA, which then inserts elsewhere among the chromosomes.

Duplication of an entire genome results in polyploidy, discussed in chapter 13. It is common in plants and some insects, but not vertebrates. (Polyploidy versus duplications can be compared to burning an entire CD versus copying only

certain songs.) If a polyploid event was followed by the loss of some genes, then peppered with additional gene duplications, the result would look much like the modern human genome (figure 16.13). The remnants of such an ancient whole-genome duplication would have become further muddled with time, as inversions and translocations altered the ancestral DNA sequence.

Ancient DNA

When comparing DNA of modern species, a researcher can easily repeat an experiment—ample samples of DNA are available directly from the sources. This isn't so for ancient DNA, such as genetic material from insects preserved in amber, which is hardened resin from pine trees. The mix of chemicals in amber entombed whatever fell into it when it was the consistency of maple syrup. Alcohols and sugars in the resin dried out the specimen, and other organic molecules acted as fixatives, keeping cellular contents in place. The resin sealed out oxygen and bacteria, preventing decomposition. Finally, other organic molecules hardened the resin over 4 to 5 million years. Today, the DNA is extracted and amplified.

Probing ancient, preserved DNA for clues to past life is romanticized in fiction. In reality, researchers at Pennsylvania State University sequenced most of the genomes in hair cells from two woolly mammoths that were flash-frozen at high altitudes in Siberia about 60,000 and 20,000 years ago. The genome of the mammoth was about 4 billion bases, compared to 3.3 million for the modern elephant, but much of the difference is due to bacteria and fungi contaminating the preserved hides. The huge animals lived on the grasslands of Siberia from 6 million years ago until about 10,000 years ago, although a few isolated populations survived until about 3,800 years ago (figure 16.14). So far mammoths appear to have very little genetic diversity, which might have made them vulnerable to disease during times of environmental stress. Comparison of the mammoth genome to the Indian and African elephant genomes may provide clues to exactly how the mammoths became extinct. Some people have suggested recreating a mammoth by inserting its unique DNA sequences into an elephant genome, and gestating a baby in an elephant.

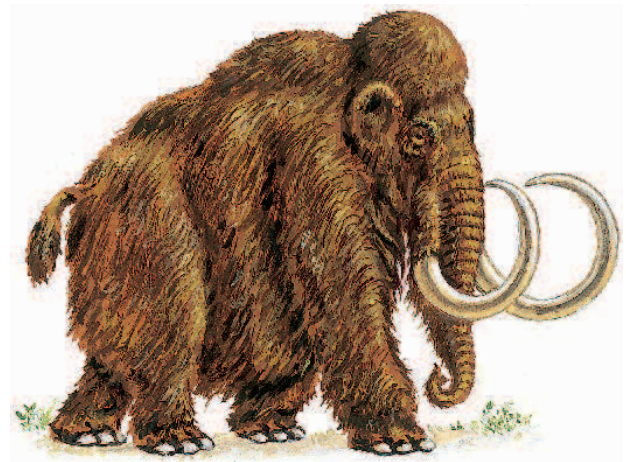


Figure 16.14 The woolly mammoth. These elephant ancestors lived perhaps as recently as 3,800 years ago. Researchers at Penn State University have sequenced nearly all of the mammoth genome, using several specimens, including two remarkably well-preserved youngsters.

Ancient DNA was first extracted in 1990, from a 17-million-year-old magnolia leaf entombed in amber. The ability to probe ancient DNA has sent researchers into the back rooms of museums, dusting off specimens of pressed leaves, pinned insects, and old bones and pelts, in search of nucleic acid clues to past life.

Comparing Chromosomes

Before gene and genome sequencing, researchers recognized that similarities in chromosome banding patterns reflect evolutionary relatedness. Human chromosome banding patterns most closely match those of chimpanzees, then gorillas, and then orangutans (table 16.2). The karyotypes of humans, chimpanzees, and apes differ from each other mostly by inversions, which occur within chromosomes.

Chromosome banding patterns are like puzzle pieces. If both copies of human chromosome 2 were broken in half,

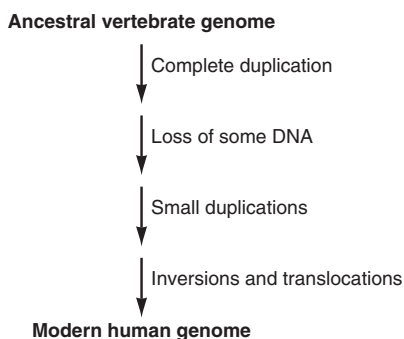


Figure 16.13 The evolution of the modern human genome.

Table 16.2	Percent of Common Chromosome Bands Between Humans and Other Species
Chimpanzees	99 ⁺ %
Gorillas	99 ⁺ %
Orangutans	99 ⁺ %
African green monkeys	95%
Domestic cats	35%
Mice	7%

we would have 48 chromosomes, as the three species of apes do, instead of 46. The banding pattern of chromosome 1 in humans, chimps, gorillas, and orangutans matches that of two small chromosomes in the African green monkey, suggesting that this monkey was ancestral to the other primates. Karyotype differences between these three primates and more primitive primates are predominantly translocations.

We can also compare chromosome patterns between species not as closely related. All mammals, for example, have identically banded X chromosomes. One section of human chromosome 1 that is alike in humans, apes, and monkeys is also remarkably similar to parts of chromosomes in cats and mice. A human even shares several chromosomal segments with a horse (figure 16.15), but our karyotype is much less like that of the aardvark, the most primitive placental mammal.

Chromosome band pattern similarities of stained material are imprecise, because a band can contain many genes that differ from those within a band at a corresponding locus in another species' genome. In contrast, DNA probes used in a FISH analysis highlight specific genes (see figure 13.9). FISH can indicate direct correspondence of gene order, or **synteny**, between species, which is solid evidence of close evolutionary relationships. For example, 11 genes are closely linked on the long arm of human chromosome 21, mouse chromosome 16, and on a chromosome called U10 in cows. However, several genes on human chromosome 3 are found near the human chromosome 21 counterpart in mice and cows. Perhaps a mammal ancestral to these three species had all of these genes together, and the genes dispersed to an additional chromosome in humans.

Comparing Proteins

Many different types of organisms use the same proteins, with only slight variations in amino acid sequence. The similarity of protein sequences is compelling evidence for descent from shared ancestors—that is, evolution. Many proteins in humans and chimps are alike in 99 percent of their amino

acids, and several are identical. When analyzing a gene's function, researchers routinely consult databases of known genes in many other organisms. Two of the most highly conserved proteins are cytochrome *c* and homeobox proteins. Another interesting conserved protein causes “double muscles.”

Cytochrome *c* is one of the most ancient and well-studied proteins. It helps to extract energy from nutrients in the mitochondria. Twenty of 104 amino acids occupy identical positions in the cytochrome *c* of all eukaryotes. The more closely related two species are, the more alike their cytochrome *c* amino acid sequence is (table 16.3). Human cytochrome *c*, for example, differs from horse cytochrome *c* by 12 amino acids, and from kangaroo cytochrome *c* by 8 amino acids. The human protein is identical to chimpanzee cytochrome *c*.

Homeobox Proteins

A class of genes that has changed little across evolutionary time is a **homeobox**, or *HOX* gene. These genes encode transcription factors that control the order in which an embryo turns on genes that ensure that anatomical parts—whether a leg, petal, or segment of a larva—develop in the right

Table 16.3 Cytochrome *c* Evolution

Organism	Number of amino acid differences from humans
Chimpanzee	0
Rhesus monkey	1
Rabbit	9
Cow	10
Pigeon	12
Bullfrog	20
Fruit fly	24
Wheat germ	37
Yeast	42

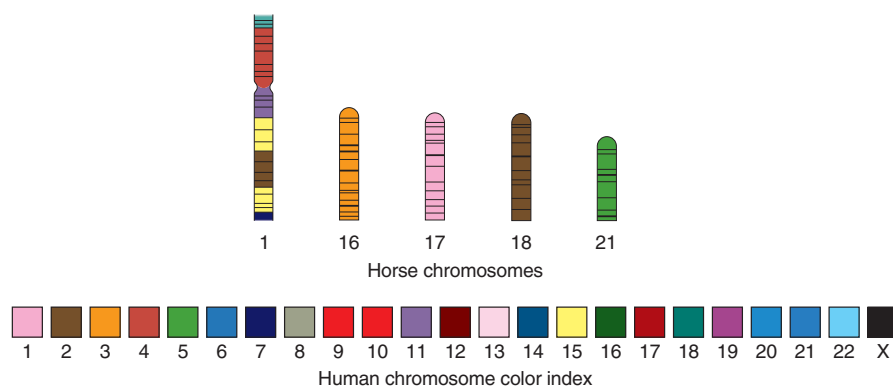


Figure 16.15 Comparing horse and human chromosomes. This color-coded display of the horse genome indicates, at a glance, regions that are highly conserved. Many horse chromosomes are very similar to human chromosomes, but note that horse chromosome 1 corresponds to different human chromosomes.

places. The highly conserved portion of a homeobox protein is a 60-amino-acid sequence. Humans and most other vertebrates have 39 *HOX* genes in four clusters. The individual genes are expressed in a sequence, in developmental time or anatomical position, that mirrors their order on the chromosome.

Mutations in the homeobox were first studied extensively in the fruit fly *Drosophila melanogaster*, mixing up body parts. *Antennapedia*, for example, has legs in place of its antennae; *proboscipedia* grows legs on its mouth. In humans, mutations in homeobox genes cause various illnesses. In a form of leukemia, a homeobox mutation shifts certain white blood cell progenitors onto the wrong developmental pathway. The misguided cells retain the rapid cell division characteristic of progenitor cells, causing the cancer. DiGeorge syndrome (MIM 188400) is also caused by a homeobox mutation. Although affected individuals hardly sprout legs from their heads, as do *Antennapedia* flies, the missing thymus and parathyroid glands and abnormal ears, nose, mouth, and throat correspond to the sites of abnormalities in the flies. **Figure 16.16** shows another human disorder caused by a mutation in a *HOX* gene.

“Double muscle” mutations vividly demonstrate how similar genes can have similar effects on different types of organisms. The double muscle phenotype is caused by lack of the protein myostatin, which normally prevents stem cells from producing muscle cells. Several members of a German family have myostatin deficiency (MIM 601788), including a boy who, at 5 years of age, could lift weights that a normal adult couldn’t, as well as several construction workers who excel in lifting boulders. Double-muscled cattle (**figure 16.17**) and chickens are prized for their extra meat, and “mighty mice” with myostatin mutations are used to study muscle overgrowth.



Figure 16.16 A human *HOX* gene mutation causes **synpolydactyly** (MIM 186000). Mutation in the *HOXD13* gene disrupts development of fingers and toes, causing a very distinctive phenotype. The third and fourth fingers are partially fused with an extra digit within the webbed material.



Figure 16.17 **Double muscling in cattle.** The mutation that causes “double muscling” is highly conserved.

Key Concepts

1. The more recently two species shared an ancestor, the more alike their DNA and protein sequences and chromosome banding patterns.
2. Chimps and humans share about 98.7 percent of protein-encoding DNA, but the proportion decreases when we consider insertions, deletions, introns, and repeats. Gene expression patterns also distinguish humans from chimps. The human genome likely duplicated during its evolution.
3. DNA from preserved extinct organisms can be amplified and compared to sequences in modern species.
4. Chromosome banding pattern similarities and amino acid sequences of highly conserved proteins reflect species relationships.

16.3 Molecular Clocks

A clock measures the passage of time by moving its hands through a certain degree of a circle in a specific and constant interval of time—a second, a minute, or an hour. Similarly, an informational molecule can be used as a “molecular clock” if its building blocks are replaced at a known and constant rate.

The similarity of nuclear DNA sequences in different species can be used to estimate the time when the organisms diverged from a common ancestor, if the rate of base substitution mutation is known. For example, many nuclear genes studied in humans and chimpanzees differ in 5 percent of their bases, and substitutions occur at a rate of 1 percent per 1 million years. Therefore, 5 million years have presumably passed since the two species diverged. Mitochondrial DNA (mtDNA) sequences may also be tracked in molecular clock studies, as we will soon see.

Timescales based on fossil evidence and molecular clocks can be superimposed on evolutionary tree diagrams constructed from DNA or protein sequence data. However, evolutionary trees can become complex when a single set of data can be arranged into many different tree configurations. A tree for seventeen mammalian species, for example, can be constructed in 10,395 different ways! The sequence in which the data are entered into tree-building computer programs influences the tree's shape, which is vital to interpreting species relationships. With new sequence information, the tree possibilities change.

Parsimony analysis is a statistical method used to identify an evolutionary tree likely to represent what really happened. An algorithm connects all evolutionary tree sequence data using the fewest possible number of mutational events to account for observed DNA base sequence differences. For the 5-base sequence in **figure 16.18**, for example, the data can be arranged into two possible tree diagrams. Because mutations are rare events, the tree that requires the fewest mutations is more likely to reflect reality.

Types of Genetic Markers

To track ancient human migration patterns, researchers use the several types of markers already discussed. These include single nucleotide polymorphisms (SNPs), short tandem repeats (STRs, or microsatellites) and other copy number variants (CNVs) (see table 5.3). Markers of mitochondrial DNA (mtDNA) trace the female lineage, and markers of Y chromosome sequences trace the male lineage. Markers also follow DNA sequences that are part of autosomes. These are called “ancestry informative markers.” Sequencing mtDNA and Y chromosome DNA sequences provides information on only some of a person's ancestors.

MtDNA is ideal for monitoring recent events because it mutates faster than DNA in the nucleus—its sequences change by 2 to 3 percent per million years. Mutations accumulate faster in mtDNA because there is no DNA repair. Another advantage of typing mtDNA is that it is more abundant than nuclear DNA because mitochondria have several copies of it, and a cell has many mitochondria. When researchers are lucky enough to find fossils or ancient humans, mtDNA is the most likely DNA to be recovered.

Most of the Y chromosome DNA sequence offers the advantage of not recombining. Crossing over, which it could only do with an X chromosome because there is no second Y, would break the linkage from the past generation and therefore make tracing relationships impossible.

SNPs (single nucleotide polymorphisms) pepper mitochondrial and Y chromosome DNA, and groups of them define long DNA stretches termed **haplogroups**. (The haplotypes used to describe linkage in chapter 5 refer to shorter DNA sequences.) Y haplogroups, at last count, were considered from “A” through “T,” with several subgroups, called subclades, indicated by alternating letters and numbers. Haplogroups and their subgroups also describe mtDNA. Populations can be classified into both mtDNA and Y chromosome groups, indicating the sources of female and male lineages, respectively. Sub-Saharan Africans, for example, have Y haplogroups E1, E2, and E3a and mtDNA haplogroups L3. Europeans, however, have Y haplogroups R, I, E3b, and J, and their mtDNA haplogroup is R, which includes three subgroups.

Before the human genome was sequenced and SNPs identified throughout the genome, and before mtDNA and Y chromosome sequences were widely used, molecular clock data reflected mutations in a handful of well-studied individual

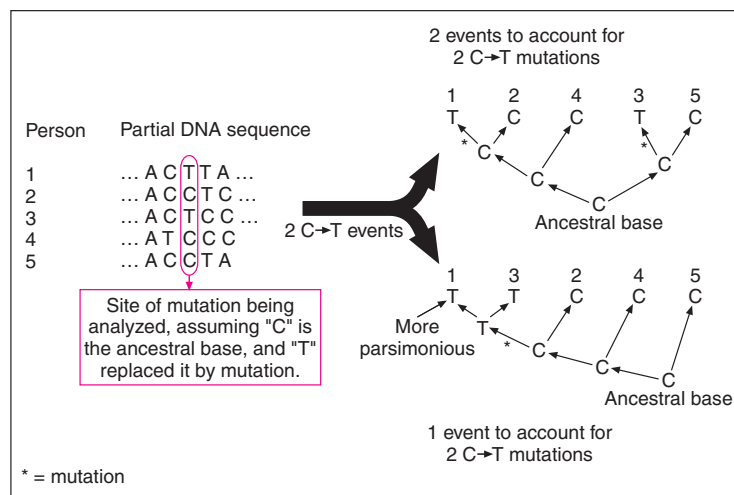


Figure 16.18 Parsimony analysis. Even a computer has trouble arranging DNA differences into an evolutionary tree showing species, population, or individual relationships. A parsimonious tree accounts for all data with the fewest number of mutations. Here, the two individuals who have a T in place of the ancestral C could have arisen in two mutational events or one, assuming that these individuals had a common ancestor. Since mutations are rare events, the more realistic scenario is one mutation.

genes. In hindsight, extrapolating from these mutation rates in individual genes to approximate the passage of time was a rather crude approach. Today, researchers use the Human Genome Diversity Panel, which is a collection of 500,000 SNPs and CNVs from more than 1,000 individuals representing 51 populations throughout the world, to see how groups of humans are connected in time and space. **Reading 16.2** discusses ancestry testing, available on the Web, that reveals individuals' mtDNA and Y chromosome haplogroups.

Neanderthals Revisited

Molecular clock data can provide clues to ancestral relationships among modern organisms and also fill gaps in the fossil record. Our increasing knowledge of the Neanderthals illustrates the recent evolution of genetic and genomic approaches to learning about the past.

Analyzing Neanderthal DNA has been difficult because it is rare and usually damaged as well as contaminated with DNA from the people handling it. But little bits of Neanderthal DNA can yield a great deal of information.

The first Neanderthal DNA sequence came from several 100-base-pair-long pieces of mtDNA that do not encode protein. These sequences mutate very rapidly because they do not affect the phenotype and are therefore not subject to natural selection. That sample, analyzed in 1997, came from a bit of arm bone from the original French Neanderthal skeleton. The DNA sequences were compared to corresponding sequences from modern *Homo sapiens*, and were found to differ at 26 positions. This is three times the number of differences between pairs of the most unrelated modern humans, and the base differences were completely different from SNPs in modern human genes. This evidence was interpreted to indicate that Neanderthals and our direct ancestors parted ways before humans left Africa, and that we did not interbreed. A small population that ended up north of the Mediterranean became the Neanderthals, and everyone else became us.

These initial results suggested that there was very little Neanderthal DNA retained in the human genome, but this analysis only considered certain mtDNA sequences. By 2006, researchers used a new DNA sequencing technology that works especially well on fragmented DNA to analyze a million bases of Neanderthal nuclear DNA. The results supported the earlier findings.

Researchers have sequenced most of the genome of a Neanderthal from a leg bone fragment found in a cave in Croatia. The man lived about 38,000 years ago; his leg bone had been sitting in a museum drawer for 30 years. While the genome is being analyzed, researchers have already discovered some interesting gene variants. The Neanderthals had variants of the *FOXP2* gene that might have enabled them to vocalize, and possibly speak. Some of them had mutations in the *MC1R* gene that gave them pale skin and red hair (**figure 16.19**).



Figure 16.19 A Neanderthal phenotype. Mutations found in Neanderthal DNA suggest that some of them may have had pale skin and red hair.

16.4 The Peopling of the Planet

Because of the scarcity of fossil evidence and limitations of extrapolating and inferring relationships from DNA sequence data, we see only glimpses of our past. However, researchers are deducing the major movements that peopled the planet (see figure 1 in Reading 16.2). Three chapters in the story of modern human origins stand out: our beginnings some 200,000 years ago; our expansion from Africa; and the populating of the New World. **Table 16.4** tells other “stories” from DNA sequence comparisons.

Mitochondrial Eve

Theoretically, if a particular sequence of mtDNA could have mutated to yield the mtDNA sequences in modern humans, then that ancestral sequence may represent a very early human or humanlike female—a mitochondrial “Eve,” or metaphorical first woman. **Figure 16.20** shows how one maternal line may have persisted.

When might this theoretical “first” woman, the most recent female ancestor common to us all, have lived? In the mid 1980s, researchers compared mtDNA sequences for protein-encoding as well as noncoding DNA regions in a variety of people, including Africans, African Americans, Europeans, New Guineans, and Australians. They concluded that the hypothesized ancestral woman lived about 200,000 years ago, in Africa. More recent analysis of mtDNA from 600 living East Africans estimated 170,000 years ago for the beginning of the modern human line, which is remarkably close to the date of the *H. sapiens idaltu* fossils. The locations of fossil evidence,

Reading 16.2

Should You Take a Genetic Ancestry Test?

About a million people so far have searched for information on their “deep ancestry” by mailing a cheek scraping or saliva sample to any of two dozen companies offering genetic genealogy services. Prices range from \$100 to \$900. These companies do not test paternity, nor do they provide health information. Instead, the goal is to assist people whose searches for the branches of their family trees using conventional documents and memories have left questions. DNA testing can answer some of them.

Most companies offer two basic types of tests: mitochondrial DNA to trace maternal lineages and Y chromosome tests to trace paternal lineages. Several markers are checked for each DNA source, and the results compared to growing databases to look for matches. “Deep ancestry” refers to assigning a match to a major part of the world—subSaharan Africa, for example—or a major population group, such as Native American or African American (**figure 1**).

DNA ancestry testing can reveal whether any two individuals living today share an ancestor, assigning an approximate generation to the “most recent common ancestor” (MRCA). The more markers tested, the more meaningful the results. If two people share all thirty-seven of thirty-seven tested markers, there is a 50 percent chance that their

MRCA was no more than two generations ago—they are so alike that other DNA variants haven’t intruded, because not enough generations have passed. Sharing twenty-five of twenty-five markers gives a 50 percent chance that the MRCA was not more than three generations ago. If they share twelve markers, there is a 50% chance that the MRCA was no longer ago than seven generations.

As is the case with direct-to-consumer genetic testing, discussed in chapter 1, ancestry testing websites offer complete and expert information—but people may still request these services with unrealistic expectations. The results are not nearly as specific as what President Barack Obama found when he visited Kenya to meet his half-siblings. But as genetic genealogy databases grow, so will the information available. Most companies inform clients when newcomers’ DNA indicates that they are related, introducing many distant cousins. Companies are also beginning to offer autosomal markers, tracing parental lines in greater detail. As a result, places of origin are becoming more specific. At first companies tested for only four parental populations—African, European, East Asian, and Native American. Now companies report on subregions of these areas.

Ancestry testing comes with some caveats:

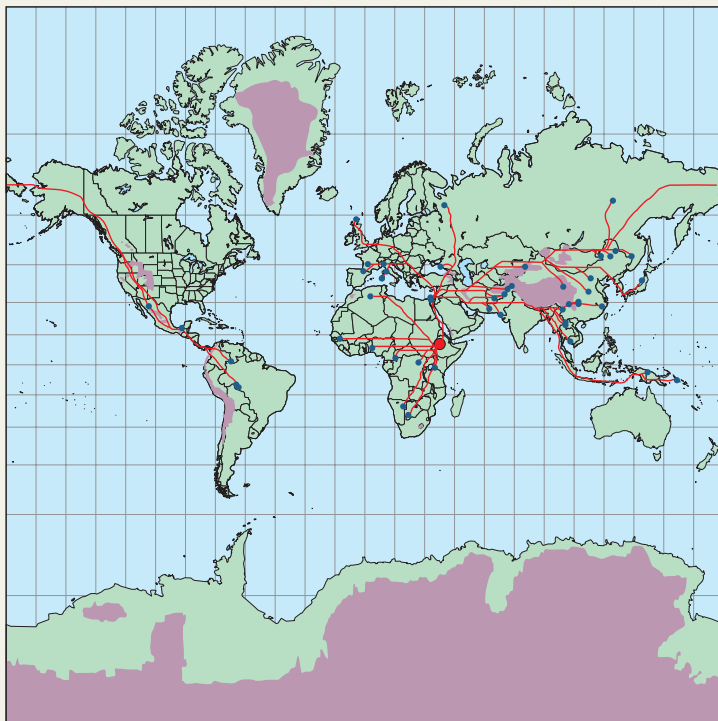


Figure 1 The peopling of the planet Humanity arose in East Africa (large red dot) and expanded to the rest of the world. This illustration depicts major migratory paths based on DNA haplogroup information, mostly from mtDNA and the Y chromosome.

- MtDNA and Y chromosome DNA testing sample considerably less than 1 percent of the genome. In those other sequences lie clues to other relatives, past and present.
- MtDNA and Y chromosome DNA trace only some lineages. You can trace your mother’s mother or your father’s father—cousins need to be tested to trace other lineages, and some lineages cannot be determined.
- Not all of the human haplogroups in the world have been discovered yet. A person’s reported geographical place of origin can change with new findings.
- A haplogroup need not come from only one geographic region, due to gene flow. People move, and so do their genes.
- Geographic regions that a haplogroup may point to may not reflect a particular ethnic or racial group, which are social, not genetic.

For people not seeking personal information but wishing to partake in a worldwide effort to map ancient migratory paths, the Genographic Project offers mtDNA or Y chromosome DNA testing for \$99. The project began with studying indigenous peoples, but then expanded to everyone. The Genographic Project is supported by National Geographic, IBM, private foundations, and academic institutions. The website is <https://www3.nationalgeographic.com/genographic/participate.html>

Table 16.4 Tales of Ancestry in Genomes

Population	DNA	History	New Findings
Tuscans (Italy)	mt	Tuscans descend from Etruria, an advanced culture predating the Roman Empire.	Unexpected Near East haplogroups found
Ashkenazi Jews	mt	Originated in 7th and 8th centuries when families from north Italy migrated to the Rhine Valley.	40% of the 8 million modern Ashkenazim descend from four women
Lebanon	Y	Modern humans arrived ~47,000 years ago, Muslims from Arabian Peninsula in 7th century CE, Christians from Crusades, 11th–13th centuries.	Current groups reflect the two religions
Island Southeast Asia (Indonesia, East Malaysia, Philippines)	mt	Modern humans arrived 50,000 years ago; rice farmers from Taiwan came 4,000 years ago and dominate linguistic and archeological evidence.	From 5,000 to 15,000 years ago, indigenous peoples expanded; Taiwanese farmer contribution was minor

such as *H. sapiens idaltu* skulls, support an African origin, and Charles Darwin suggested it, too.

One way to reach this time estimate is by comparing how much the mtDNA sequence differs among modern humans to how much it differs between humans and chimpanzees. The differences in mtDNA sequences among contemporary humans amount to 1/25 the difference between humans and chimpanzees. The two species diverged about 5 million years ago, according to extrapolation from fossil and molecular evidence. Multiplying 1/25 by 5 million gives a value of 200,000 years ago, assuming that the mtDNA mutation rate is constant over time.

Where did Eve live? Studies comparing mitochondrial and nuclear DNA sequences consistently find that Africans have the most numerous and diverse mutations. For this to be so, Africans must have existed longer than other modern

peoples, because it takes time for mutations to accumulate. In many evolutionary trees constructed by computer parsimony analysis, the individuals whose DNA sequences form the bases are from Africa. That is, other modern human populations all have at least part of an ancestral African genome, plus mutations that occurred after their ancestors left Africa.

Expansion Out of Africa

The idea of mitochondrial Eve is part of the “out of Africa” view, or replacement hypothesis, of human origins. It states that about 200,000 years ago, *H. sapiens* evolved from an *H. erectus* or other *Homo* population in Africa. This may have occurred quickly, in small, isolated pockets, or gradually across a broader swath of the continent. However it happened, eventually descendants of some of these early *H. sapiens* expanded

out of Africa about 56,000 years ago. Some remained. An alternate view, the multiregional hypothesis, has been largely disproved. It maintained that modern humans arose in several places, gradually emerging on a global scale in which people from all over mixed—not the isolated pockets of peoples that the replacement hypothesis envisions.

Data from mtDNA, Y chromosome DNA, and markers on the autosomes indicate that the peopling of the world was a series of founder effects as groups left Africa, perhaps when the Sahara desert periodically grew wetter. These movements yielded “chains of colonies” that may have overlapped and merged when neighbors met, and genes flowed from one region to another. Geographic and climatic barriers periodically shrank human populations,

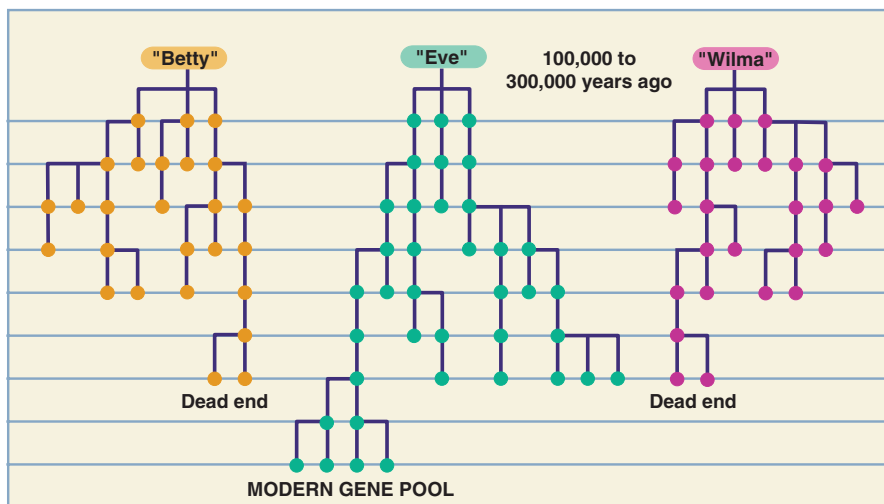


Figure 16.20 Mitochondrial Eve. According to the mitochondrial Eve hypothesis, modern mtDNA retains some sequences from a figurative first woman, “Eve,” who lived in Africa 300,000 to 100,000 years ago. In this schematic illustration, the lines represent generations, and the circles, females. Lineages cease whenever a woman does not have a daughter to pass on the mtDNA.

while natural selection and genetic drift narrowed the African gene pool. At the same time, new mutations established the haplogroups that are consulted to trace ancestry.

Many questions remain. When did groups leave Africa? Did they meet and mate? How and when did population bottlenecks carve out modern gene pools? Were Neanderthals the only offshoots, known because we happened to find their fossils? Probing genes and genomes for clues to the past is like a photo slowly coming into focus, capturing a portrait of our origins.

Populating the New World

People spread across Eurasia by 40,000 years ago, as well as elsewhere, and lastly through Siberia (**figure 16.21**). From here they could cross the Bering Land Bridge, which emerged between Siberia and Alaska during times when the glaciers had retreated. The land bridge stretched for about 1,000 miles from north to south, emerging as winds from the southwest kept snow away. The areas for several hundred miles on either side of the bridge, and the bridge itself, are called Beringia.

Sometime between 23,000 and 19,000 years ago, a severe population bottleneck affected the people in Beringia. Only about 1,000 of them survived the journey over the bridge from Siberia, and some of them continued southward along the Pacific coastline. As the ice age ended about 18,000 years ago, the population began a period of rapid expansion that lasted 3,000 years. They spread through the Americas as the founding Native Americans. The mitochondrial DNA evidence that paints this picture is particularly valuable because the coastal migratory path is now underwater, hiding archeological clues.

Today, Native Americans carry a very distinctive genetic nametag that reflects the long-ago trek across the land bridge: five mtDNA haplogroups (A, B, C, D and X) and two Y chromosome haplogroups (C and Q). These markers are seen in all indigenous populations in southern Siberia too, indicating a single gene pool traveling in a single migration. DNA sequence information extrapolated back from present-day Native American populations is consistent with molecular clock data from ancient DNA—telling a wonderful scientific story. In addition, Native American populations have a STR marker not seen

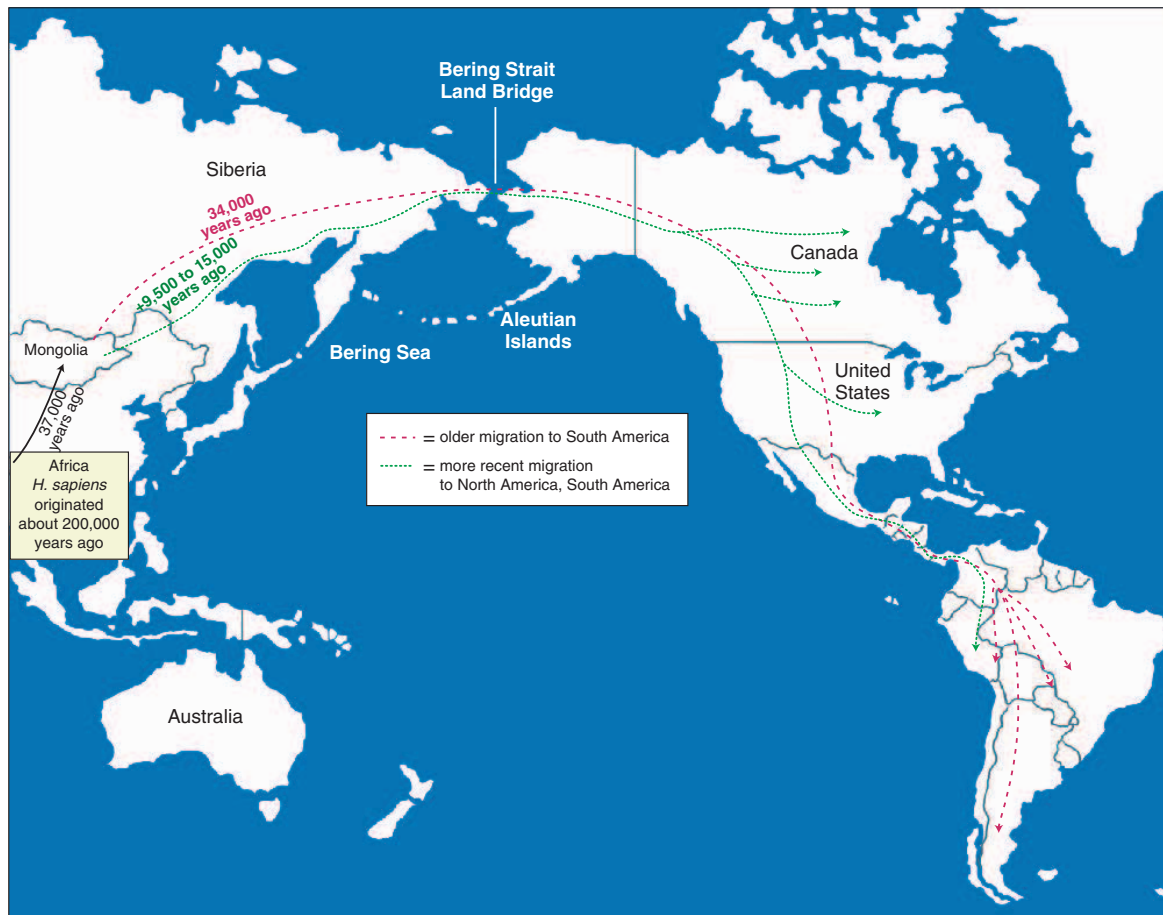


Figure 16.21 Tracing human origins. Analyses of mitochondrial DNA and Y chromosome DNA sequences reveal that the ancestors of Native Americans probably came from Mongolia in one migration.

in eastern Siberian peoples, indicating a genetic change that occurred after the crossing.

A comparison of 678 autosomal STR markers from 29 Native American populations and 49 other indigenous groups worldwide found that Native Americans are very different from other populations, yet are very much like each other—as might be expected from a multigenerational journey southward by a small but hardy group, along the coastline of the Americas.

By 14,000 years ago, Native Americans had arrived inland. Evidence comes from their DNA in coprolites—fossilized excrement—discovered in Oregon, Wisconsin, and even Florida. Genetic evidence also suggests that some Native American populations died out, perhaps descendants of those who chose to stay in the harsh north of the new land. A clump of hair from a Paleo-Eskimo found in Greenland dates from 4,000 years ago. The Paleo-Eskimos migrated into Alaska, Canada, and Greenland about 4,500 years ago. Surprisingly, his mtDNA haplogroup was neither Native American nor modern Eskimo. The conclusion: The earliest Eskimos came from a different migration than the one that was ancestral to Native

Americans and modern Eskimos, and left no present-day descendants. The discovery of the 4,000-year-old non-Native American/non-Eskimo reveals a major limitation of genetic anthropology: We cannot fit in the puzzle pieces that we cannot find, or that no longer echo in modern gene pools.

Key Concepts

1. Molecular clocks apply mutation rates to timescales to estimate when two individuals or types of organisms most recently shared ancestors.
2. Different genes evolve at different rates. Parsimony analysis selects likely evolutionary trees from DNA data.
3. Mitochondrial DNA clocks trace maternal lineages, and Y chromosome sequences trace paternal lineages.
4. Molecular clocks have been used to examine the origin and migrations of modern humans.

Summary

16.1 Human Origins

1. The first primates were rodentlike insectivores that lived about 60 million years ago. By 30 to 40 million years ago, monkeylike *Aegyptopithecus* lived. **Hominoids**, ancestral to apes and humans, lived 22 to 32 million years ago.
2. **Hominins**, ancestral to humans only, appeared about 19 million years ago. They were more upright, dwelled on the plains, and had smaller brains than their ancestors.
3. At least three types of hominins lived about 6 million years ago, shortly after the split from the chimp lineage.
4. The australopithecines preceded and then coexisted with *Homo habilis*, who lived in caves, had strong family units, and used tools extensively. *Homo erectus* was a contemporary who outlasted *H. habilis*, lived in societies, and used fire. *Homo sapiens idaltu* lived about 156,000 years ago, and looked like us.
5. Neanderthals split from the human lineage about 516,000 years ago but died out. Modern humans appeared about 200,000 years ago.

16.2 Molecular Evolution

6. Molecular evolution considers differences at the genome, chromosome, protein, or DNA sequence levels with mutation rates to estimate species relatedness.
7. Humans and chimps share 98.7 percent of their protein-encoding gene sequences. Indels, introns, and repeats create genome differences between humans and chimps, which also differ in gene expression.
8. The human genome shows many signs of past duplication.
9. Amplifying ancient DNA is difficult because contamination may occur.

10. Closely related species have similar chromosome banding patterns. Genes in the same order on chromosomes in different species are **syntenic**.
11. For a highly conserved gene or protein, DNA sequence is similar or identical in different species, and presumably indicates importance and descent from common ancestors.

16.3 Molecular Clocks

12. Gene sequence information from several species may be used to construct evolutionary tree diagrams, and a molecular clock based on the known mutation rate of the gene applied. Different genes mutate at different rates. Molecular trees indicate when species diverged from shared ancestors.
13. Parsimony analysis selects the evolutionary trees requiring the fewest mutations, which are therefore the most likely.
14. Molecular clocks based on mtDNA date recent events through the maternal line because this DNA mutates faster than nuclear DNA. Y chromosome genes trace paternal lineage. Markers (SNPs, STRs, and CNVs) in both types of evidence are used to study human origins and expansions. Groups of linked markers inherited together form **haplogroups**.

16.4 The Peopling of the Planet

15. The rate of mtDNA mutation and current mtDNA diversity can be extrapolated to hypothesize that a theoretical first woman lived, in Africa, about 200,000 years ago. People began to leave Africa about 56,000 years ago.
16. The peopling of the planet was a series of migrations and founder effects, with genetic diversity decreasing from that of the ancestral African population, but new mutations occurring.
17. The peopling of the Americas occurred after the last ice age, permitting people to cross the Bering Strait from Siberia.

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Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



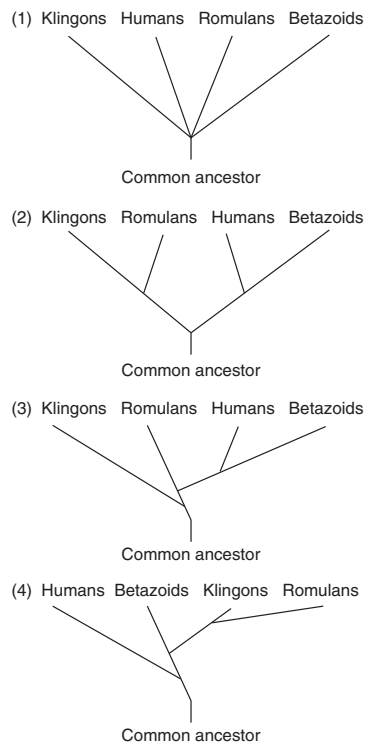
Review Questions

- Arrange the following primates in the order in which they lived, indicating any that may have overlapped in time.
 - Homo erectus*
 - Australopithecus anamensis*
 - Dryopithecus*
 - Neanderthals
 - Ardipithecus*
 - Homo habilis*
 - Australopithecus garhi*
 - Homo sapiens idaltu*
- What is the difference between a hominoid and a hominin?
- Some anthropologists classify chimpanzees along with humans in genus *Homo*. How does this conflict with fossil evidence of the *Australopithecus* species? With DNA sequence evidence from Neanderthals?
- Give an example of how a single gene difference can have a profound effect on the phenotypes of two species.
- Give an example of molecular evidence that is consistent with fossil or other evidence.
- Describe the type of information that Y chromosome and mitochondrial DNA sequences provide.
- List three aspects of development, anatomy, or physiology that were important in human evolution.
- Why is a DNA sequence that is highly conserved among humans and chimps, gorillas, and orangutans unlikely to vary greatly among human populations?
- Explain how indels could cause the divergence of our genome sequence from that of chimpanzees, yet not contribute to observable differences between the two species.
- Cite two ways that humans and chimps can differ greatly at the genetic level, but still be very alike in terms of DNA sequence.
- What types of information are needed to construct an evolutionary tree diagram? What assumptions are necessary? What are the limitations of these diagrams?
- Researchers compare a number of types of information in the human and chimp genomes, including SNPs, CNVs, STRs, indels, and linkage patterns, discussed in this and other chapters. Define each of these types of information.
- How can the human and chimp genomes be nearly 99 percent alike in DNA sequence, yet still be different?

Applied Questions

- Select an example from this chapter and explain how it illustrates one of the forces of evolutionary change discussed in chapter 15 (natural selection, nonrandom mating, migration, genetic drift, or mutation.)
- Create a narrative of how a group of australopithecines might have branched off into *Homo*.
- Why did many species of *Australopithecus* live at one time, but only one of *Homo* lives today?
- A geneticist aboard a federation starship must deduce how closely related Humans, Klingons, Romulans, and Betazoids are. Each organism walks on two legs, lives in complex societies, uses tools and technologies, looks similar, and reproduces in the same manner. Each can interbreed with any of the others. The geneticist finds the following data:
 - Klingons and Romulans each have 44 chromosomes. Humans and Betazoids have 46 chromosomes. Human chromosomes 15 and 17 resemble part of the same large chromosome in Klingons and Romulans.
 - Humans and Klingons have 97 percent of their chromosome bands in common. Humans and Romulans have 98 percent of their chromosome bands in common, and Humans and Betazoids show 100 percent correspondence. Humans and Betazoids differ only by an extra segment on chromosome 11, which appears to be a duplication.
 - The cytochrome *c* amino acid sequence is identical in Humans and Betazoids, differs by one amino acid between Humans and Romulans, and differs by two amino acids between Humans and Klingons.
 - The gene for collagen contains 50 introns in Humans, 50 introns in Betazoids, 62 introns in Romulans, and 74 introns in Klingons.
 - Mitochondrial DNA analysis reveals many more individual differences between Klingons and Romulans than between Humans and Betazoids.

- Suggest a series of chromosomal abnormalities or variants that might explain the karyotypic differences among these four types of organisms.
- Which are our closest relatives among the Klingons, Romulans, and Betazoids? What is the evidence for this?
- Are Klingons, Romulans, Humans, and Betazoids distinct species? What information reveals this?
- Which of the evolutionary tree diagrams is consistent with the data?



- Why might it be important to identify DNA sequences that chimps have that humans do not, as well as identify sequences unique to humans?
- Several women have offered to be inseminated with sperm from the Ice Man, who died 5,300 years ago in the Alps. If sperm could have been recovered, and a woman inseminated, what do you think the child would be like?
- Do you think it is a good idea to attempt to clone a woolly mammoth by inserting its genome into an elephant's egg? Cite a reason for your answer.
- A man with white skin routinely checks the box on forms about personal information for "African American," claiming that we are all, if we go back far enough in our family trees, from Africa. Is he correct?
- Can ancestry testing help to dispel race-based problems, or worsen them? Cite a reason for your answer.
- Hatshepsut was a pharaoh who ruled Egypt more than 3,000 years ago. Although she led many successful projects and the economy soared during her two-decade reign, she had been erased from written records. In 2008, her mummy was discovered and tentatively identified from a distinctive tooth. DNA analysis confirmed the finding, but Egypt did not allow removal of archeological samples. So, with the help of a supplier of DNA sequencing machines, the technology came to the museum. What information was needed to confirm that the mummy was, indeed, Hatshepsut, part of the royal line?
- In Central Africa, the Mbuti Pygmies are hunter-gatherers who live amid agricultural communities of peoples called the Alur, Hema, and Nande. Researchers compared autosomal STRs, mitochondrial DNA, and Y chromosome DNA haplogroups among these four types of people. The pygmies had the most diverse Y chromosomes, including about a third of the sequence the same as those among the agricultural groups, who had greater mtDNA diversity than the pygmies. None of the agricultural males had pygmy Y DNA sequences. Create a narrative of gene flow to explain these findings.

Web Activities

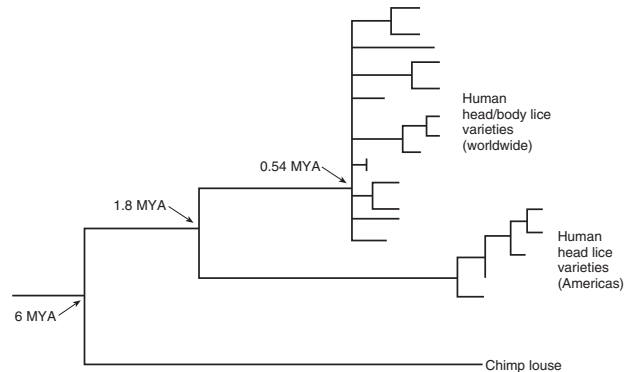
- Go to <http://www.peoplesoftheworld.org/> or a similar website. Select an indigenous people, do further research, and describe their habitat and culture, and any distinctive health strengths or problems. To what extent do you think genetics is responsible for the state of their health? Explain your reasoning.
- Consult the website for the Genographic Project, and read the material under "Atlas of Human Journey."
 - What does "deep ancestry" mean?
 - Explain why analyzing your mtDNA or Y chromosome DNA cannot provide a complete picture of your ancestry.
 - Explain how a female can trace her paternal lineage if she doesn't have a Y chromosome.
 - Would you want your ancestry information and identity posted on the Genographic database so that current cousins can contact you?

Case Studies and Research Results

14. Consider the following brain sizes:

Animal	Brain volume (cubic centimeters)
<i>Homo sapiens sapiens</i> (us)	1400–1500
<i>Homo sapiens idaltu</i>	1450
<i>Homo erectus</i>	1000–1250
<i>Australopithecus</i>	380
Chimpanzee	350–380
<i>Homo floresiensis</i>	380
Gorilla	350–450

- Explain how these data either support or refute the hypothesis that increasing brain size correlates to increased intelligence.
 - Explain a limitation of the ways in which we learn about our ancestors or relatives that might account for confusing results when we compare traits.
15. Austrian researchers discovered at an ancient burial site in the eastern part of their nation remains of two newborns, their preserved skeletons facing each other in a fetal position. The bones were small and thin, the heads small. The remains were covered with red clay, in which ornaments were embedded.
- What was the likely species of the newborns?
 - Describe two other types of information that might explain a little about the culture of the hominins represented in the remains.
16. For more than 20 million years, lice have lived on the skins of primates. Researchers compared a 1,525-base-pair sequence of mtDNA among modern varieties of lice, and, applying the mutation rate, derived the evolutionary tree in the next column. It depicts a split in the louse lineage, with one group of head and body lice living throughout the world, and another group of only head lice living in the Americas.



- What events in human evolution roughly correspond to the branch points in the louse evolutionary tree?
 - What might be the significance of the similarity between the evolutionary trees for lice and humans?
 - The researchers interpreted their findings to indicate that lice moved from archaic humans to modern humans. What is the evidence for this hypothesis? What other types of evidence or background information might make it more convincing?
 - What is a limitation of this research?
17. A Y chromosome haplotype consists of specific mutations for the *SRY* gene and genes called *M96* and *P29*. Among modern Africans, there are three variants of this haplotype. Two of them are found only among Africans, but the third variant of the Y haplotype, called E3, is also seen in western Asia and in parts of Europe. Researchers examined specific subhaplotypes (variations of the variations) and found that one type, called E-M81, accounts for 80 percent of the Y chromosomes sampled in northwest Africa, diminishing sharply in incidence to the east, and not present in sub-Saharan Africa at all. That same haplotype is found in a small but significant percentage of the Y chromosomes in Spain and Portugal. Consult a map, and propose a scenario for this gene flow. What further information would be useful in reconstructing migration patterns?

Forensics Focus

18. In many African cultures, “family” is not dictated by genetics, but by who cares for whom. Any adult can be “mother” or “father” to any child. Since the early 1990s, many parts of East Africa have been under civil war. Thousands of Africans have asked to be admitted to the United States to join relatives. The “family reunification resettlement program” enables parents, siblings, and children of U.S. citizens to come to the U.S. In 2008, addressing rumors that many people were lying that they were related to people in the United States, the State Department began to ask refugees from Kenya to voluntarily provide a DNA sample, to be compared to that of the U.S. citizen claimed to be a relative. When the DNA testing turned

up many cases of people claiming to be family who were not blood relatives, the resettlement program was stopped.

- Do you think that DNA testing should have been imposed on people seeking asylum in the United States from their war-torn countries?
- Should the people have been compelled to have their DNA tested?
- How should cultural definitions, such as that of “family,” be handled?
- How should the situation be resolved?

PART 5 Immunity and Cancer



Rheumatoid arthritis causes great deformity of the hands and very painful joints. It isn't usually inherited, but is associated with altered expression of certain genes.

CHAPTER

17

Genetics of Immunity

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Gene Expression In Rheumatoid Arthritis

Imagine not being able to hold a pen, or turn a doorknob. For someone with rheumatoid arthritis (RA), simple tasks become impossible. The joints become chronically inflamed, causing painful “flares.” Great deformity results.

RA is a classic autoimmune disease, in which the immune system attacks the tissues. RA does not follow a Mendelian pattern of inheritance in families, and is not more likely to occur in both members of MZ (identical) twins compared to both members of DZ (fraternal) twins. However, MZ twins in whom one has RA and the other doesn't provide a way to investigate the nongenetic factors that contribute to the disease.

Researchers withdrew fluid from the joints of eleven pairs of MZ twins in whom one had RA and the other didn't. They extracted DNA from cells in the fluid and applied it to DNA microarrays that detected expression (mRNA production) of 20,000 genes. Three genes were greatly overexpressed in the twins who had RA, and the corresponding proteins were overabundant in their joint fluid.

All three proteins had never been associated with RA, but they made sense. One destroys bone and cartilage. Another deactivates the hormone cortisol, whose levels are diminished in RA. The third protein stimulates blood vessel formation, which enhances inflammation. Researchers are developing drugs that target the three proteins, and investigating environmental influences that may have triggered RA in one identical twin but not the other.

17.1 The Importance of Cell Surfaces

We share the planet with plants, microbes, fungi, and other animals, but can become ill when they, or their parts, enter our bodies. The human immune system protects us against this happening. It is a mobile army of about 2 trillion cells, the biochemicals they release, and the organs where they are produced and stored.

Protection against infection is based on the ability of the immune system to recognize “foreign” or “nonself” cell surfaces that are not part of the body. These include surfaces of microorganisms such as bacteria and yeast; nonliving “infectious agents” such as viruses and prions; and even some tumor cells and transplanted cells. Then, the immune system carries out a highly coordinated attack that includes both general and highly specific responses. Organisms or infectious agents that cause disease are called pathogens. **Reading 17.1** highlights one common type of pathogen—viruses. **Figure 17.1** shows another—bacteria.

Understanding how genes control immunity makes it possible to enhance or redirect the system’s ability to fight disease. Mutations can impair immune function, causing immune deficiencies, autoimmune disorders, allergies, and cancer. Genes affect immunity by conferring susceptibilities or resistances to certain infectious diseases. Like other inherited characteristics, degree of immune protection varies from person to person. This may be why one person suffers frequent respiratory infections whereas another is rarely ill. A recent study of the immune systems of people who survived the 1918 flu pandemic, which killed 50 million people, revealed that their antibodies—to this day—can still rapidly destroy a flu virus. The elderly flu survivors enjoy another aspect of the immune response—it remembers.

A few types of single genes encode proteins that exert powerful effects on immunity. **Antibodies** and **cytokines** are

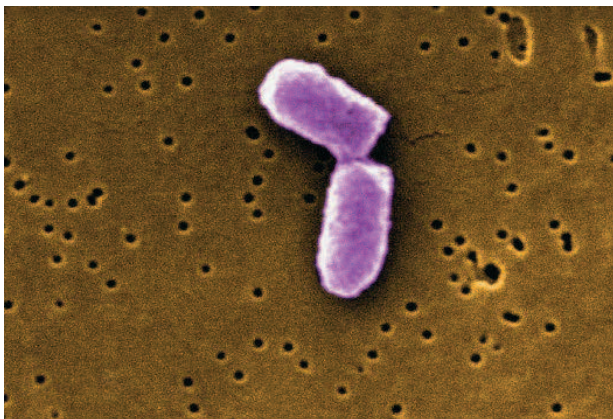


Figure 17.1 A bacterial pathogen. *Escherichia coli* is a normal resident of the human small intestine, but under certain conditions can produce a toxin that causes severe diarrhea (“food poisoning”) and can damage the kidneys.

proteins that directly attack foreign antigens. An **antigen** is any molecule that can elicit an immune response. Antigens are typically proteins or carbohydrates. Genes also specify the cell surface antigens that mark the body’s cells as “self.”

Blood Groups

Some of the antigens that dot our cell surfaces determine blood types. Figures 5.3 and 5.4 describe the familiar ABO blood types. We actually have twenty-nine major blood types based on protein and carbohydrate antigens on the surfaces of red blood cells. Each of these types includes many subtypes, generating hundreds of ways that the topographies of our red blood cells differ from individual to individual. **Table 17.1** lists a few blood groups.

For blood transfusions, blood is typed and matched from donor to recipient. For more than a century, an approach called serology typed blood according to red blood cell antigens. A newer way to type blood is to identify the *instructions* for the cell-surface antigens—that is, the genes that encode these proteins. This approach, termed genotyping, is being tested in Europe and Canada using a tiny device called a BLOOD-chip that detects 100 distinct DNA “signatures.” Genotyping is especially useful for people who have a chronic disorder that requires multiple transfusions, such as leukemia or sickle cell disease. They produce so many antibodies against so many types of donor blood that it is often difficult to determine their blood types by serology. DNA typing, however, can tell.

The Major Histocompatibility Complex

Many of the proteins on our cell surfaces are encoded by genes that are part of a 6-million-base-long cluster on the short arm of chromosome 6 called the **major histocompatibility complex** (MHC). The MHC includes about 70 genes, and about 50 percent of the genetic influence on immunity stems from these genes. MHC genes are classified into three functional groups. Class III

Table 17.1 Blood Groups

Blood Group (MIM)	Description
MN (111300)	Codominant alleles <i>M</i> , <i>N</i> , and <i>S</i> determine six genotypes and phenotypes. The antigens bind two glycoproteins.
Lewis (111100)	Allele <i>Le</i> encodes fucosyltransferase (FUT3) that adds an antigen to the sugar fucose, which the product of the <i>H</i> gene places on red blood cells. <i>H</i> gene expression is necessary for the ABO phenotype (see section 5.1). People with <i>LeLe</i> or <i>Lele</i> have the Lewis antigen on red blood cells and in saliva. People of genotype <i>lele</i> do not.
Secretor (182100)	People with <i>Se</i> allele secrete A, B, and H antigen in body fluids.

Reading 17.1

Viruses

We all know how miserable a viral infection can make us feel. Many of the aches and pains that accompany a viral infection are in fact, actions of the immune system.

A virus is a single or double strand of RNA or DNA wrapped in a protein coat, and in some types, an outer envelope, too. A virus can reproduce only if it enters and uses a host cell's energy resources, protein synthetic machinery, and secretion pathway. It is a stunningly streamlined structure. A virus may have only a few protein-encoding genes, but many copies of the same protein can assemble to form an intricate covering, like the panes of glass in a greenhouse. Ebola virus, for example, has only seven types of proteins, but they assemble into a structure capable of reducing a human body to little more than a bag of blood and decomposed tissue (**figure 1**). In contrast, the smallpox virus has more than 100 different types of proteins, and HIV is also complex.

Viruses are with us all the time—not only when we are ill. Part of the DNA sequence of some human chromosomes includes viral DNA sequences that are vestiges of past infections, perhaps passed, silently, from distant ancestors. Many DNA viruses reproduce by inserting their DNA into the host cell's DNA. In contrast, an RNA virus must first copy its RNA into DNA before it can insert into a human chromosome. A viral enzyme called reverse transcriptase does this. Certain RNA viruses are called retroviruses because they transmit genetic information opposite the usual direction—instead of DNA to RNA to protein, viral RNA is copied into DNA, which may then be copied back into RNA to guide the synthesis of viral proteins. HIV is a retrovirus.

Once viral DNA integrates into the host cell's DNA, it can either remain and replicate along with the host's DNA without causing harm,

or it can take over and kill the cell. Activated viral genes direct the host cell to replicate viral DNA and then use it to manufacture viral proteins. The cell bursts, releasing many new virus copies into the body.

Diverse viruses infect all types of organisms. Their genetic material cannot repair itself, so the mutation rate may be high—which is one reason why we cannot develop an effective vaccine against HIV or the common cold, and why new influenza vaccines must be developed each year.

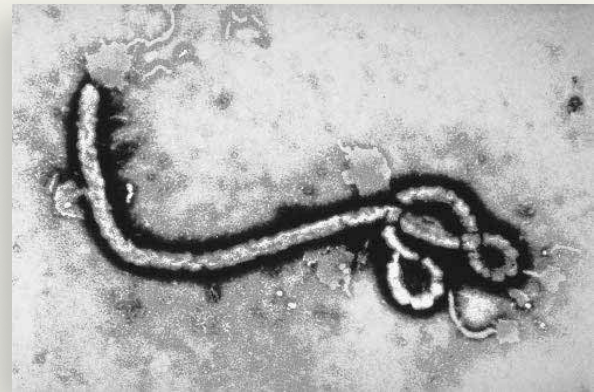


Figure 1 Ebola virus is a single strand of RNA and seven proteins. People become infected when they touch body fluids of an infected individual.

MHC genes encode proteins that are in blood plasma (the liquid portion of blood, discussed in section 11.1) and that provide nonspecific immune functions. Class I and II genes of the MHC encode the **human leukocyte antigens (HLA)**, first studied in leukocytes (white blood cells). The HLA proteins link to sugars, forming branchlike glycoproteins that extend from cell surfaces. Some HLA glycoproteins latch onto bacterial and viral proteins, displaying them like badges to alert other immune system cells. This action, called antigen processing, is often the first step in an immune response. The cell that displays the foreign antigen is called an **antigen-presenting cell**. **Figure 17.2** shows how a large cell called a macrophage displays bacterial antigens. Certain white blood cells called T cells (or T lymphocytes) are also antigen-presenting cells. Class I and II HLA proteins differ in the types of immune system cells they alert.

HLA antigens identify all cells of a person as “self,” or belonging to the same individual. In addition to these common HLA markers are more specific markers that distinguish particular tissue types. Class I includes three genes that vary greatly and are found on all cell types, and three other genes

that are more restricted in their distribution. Class II includes three major genes whose encoded proteins are found mostly on antigen-presenting cells.

Individuals have an overall HLA “type” based on the six major HLA genes. Only 2 in every 20,000 unrelated people match for the six major HLA genes by chance. When transplant physicians attempt to match donor tissue to a potential recipient, they determine how alike the two individuals are for these six genes. Usually at least four of the genes must match for a transplant to have a reasonable chance of success. Before DNA profiling, HLA typing was the predominant type of blood test used in forensic and paternity cases to rule out involvement of certain individuals. However, HLA genotyping has become very complex because hundreds of alleles are now known.

A few disorders are very strongly associated with inheriting particular HLA types. This is the case for ankylosing spondylitis, which inflames and deforms vertebrae. A person with either of two particular subtypes of an HLA antigen called B27 is 100 times as likely to develop the condition as someone who lacks either form of the antigen. HLA-associated risks are not absolute. More than 90 percent of people who suffer from

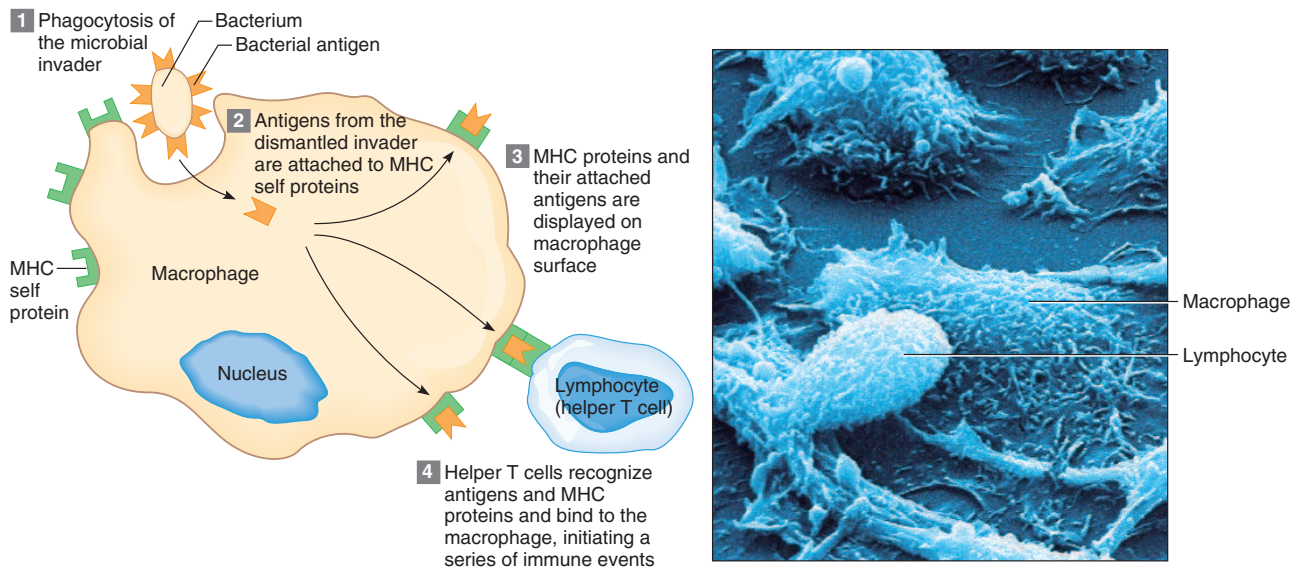


Figure 17.2 Macrophages are antigen-presenting cells. A macrophage engulfs a bacterium, then displays foreign antigens on its surface, which are held in place by major histocompatibility complex (MHC) self proteins. This event sets into motion many immune reactions.

ankylosing spondylitis have the B27 antigen, which occurs in only 5 percent of the general population. However, 10 percent of people who have ankylosing spondylitis do *not* have the B27 antigen, and some people who have the antigen never develop the disease.

Key Concepts

1. The immune system consists of cells and biochemicals that distinguish self from nonself antigens.
2. Pathogens include microorganisms and infectious agents.
3. Blood types result from self antigen patterns on red blood cells. HLA cell surface proteins establish self and display foreign antigens.

17.2 The Human Immune System

The immune system is a network of vessels called lymphatics that transport lymph fluid to bean-shaped structures throughout the body called lymph nodes. The spleen and thymus gland are also part of the immune system (**figure 17.3**).

Lymph fluid carries white blood cells called lymphocytes and the wandering, scavenging macrophages that capture and degrade bacteria, viruses, and cellular debris. **B cells** and **T cells** are the two major types of lymphocytes.

The genetic connection to immunity is the proteins required to carry out an immune response. The immune response attacks

pathogens, cancer cells, and transplanted cells with two lines of defense—an immediate generalized **innate immunity**, and a more specific, slower **adaptive immunity**. These defenses act after various physical barriers block pathogens. **Figure 17.4** summarizes the basic components of the immune system, discussed in the following sections.

Physical Barriers and Innate Immunity

Several familiar structures and fluids keep pathogens from entering the body in the innate immune response: unbroken skin, mucous membranes such as the lining inside the mouth, earwax, and waving cilia that push debris and pathogens up and out of the respiratory tract. Most microbes that reach the stomach perish in a vat of churning acid or are flushed out in diarrhea. These physical barriers are nonspecific—that is, they keep out anything foreign, not just particular pathogens.

If a pathogen breaches these physical barriers, innate immunity provides a rapid, broad defense. The term *innate* refers to the fact that these general defenses are in the body, ready to function should infection threaten. A central part of the innate immune response is **inflammation**, a process that creates a hostile environment for certain types of pathogens at an injury site. Inflammation sends in cells that engulf and destroy pathogens. Such cells are called phagocytes, and their engulfing action is phagocytosis (**figure 17.5**). Certain types of white blood cells and the large, wandering macrophages are phagocytes. Also at the infection site, plasma accumulates, which dilutes toxins and brings in antimicrobial chemicals. Increased blood flow with inflammation warms the area, turning it swollen and red.

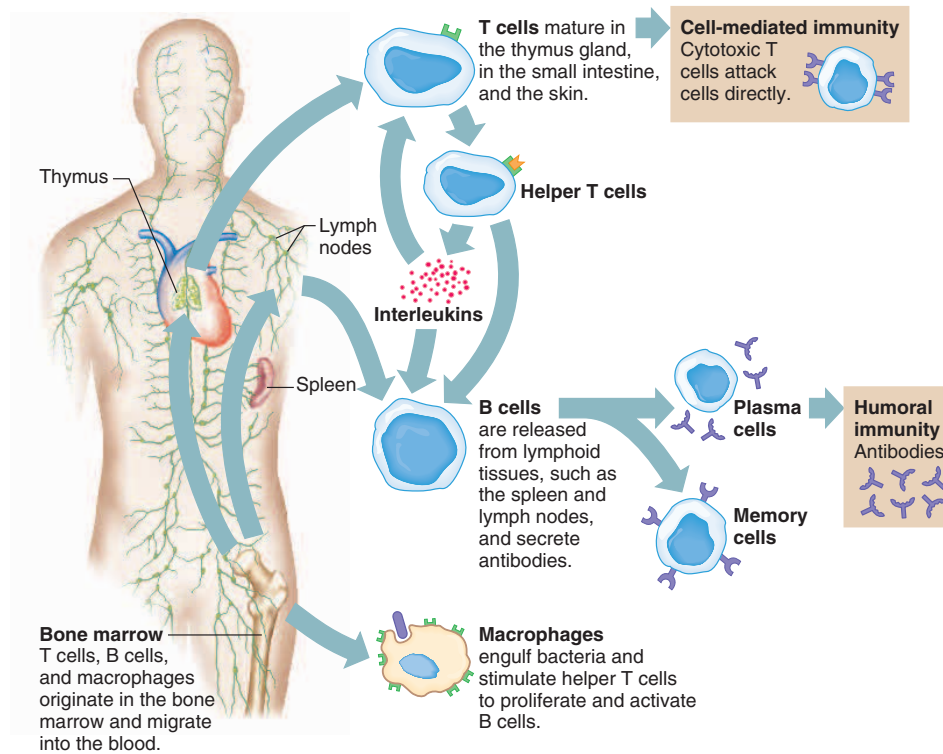


Figure 17.3 The immune system produces diverse cells. T cells, B cells, and macrophages build an overall immune response. All three types of cells originate in the bone marrow and circulate in the blood.

In addition to inflammation, three classes of proteins participate in innate immunity. These are the complement system, collectins, and cytokines. Mutations in the genes that encode these proteins increase susceptibility to infection.

The **complement system** consists of plasma proteins that assist, or complement, several other defenses. Some complement proteins puncture bacterial plasma membranes, bursting the cells. Others dismantle viruses or trigger release of histamine from mast cells, another type of immune system cell that is involved in allergies. Histamine dilates blood vessels, sending fluid to the infected or injured area. Still other complement proteins attract phagocytes to an injury site.

Collectins broadly protect against bacteria, yeasts, and some viruses by detecting slight differences in their surfaces from human cells. Groups of human collectins correspond to the surfaces of different pathogens, such as the distinctive sugars on yeast, the linked sugars and lipids of certain bacteria, and the surface features of some viruses.

Cytokines play roles in both innate and adaptive immunity. As part of the innate immune response, cytokines called **interferons** alert other components of the immune system to the presence of cells infected with viruses. These cells are then destroyed, which limits the spread of infection.

Interleukins are cytokines that cause fever, temporarily triggering a higher body temperature that directly kills some infecting bacteria and viruses. Fever also counters microbial

growth indirectly, because higher body temperature reduces the iron level in the blood. Bacteria and fungi require more iron as the body temperature rises; a fever-ridden body stops their growth. Phagocytes also attack more vigorously when the temperature rises. Tumor necrosis factor is another type of cytokine that activates other protective biochemicals, destroys certain bacterial toxins, and attacks cancer cells. Many of the aches and pains we experience from an infection are actually due to the immune response, not directly to the actions of the pathogens.

Adaptive Immunity

Adaptive immunity must be stimulated into action. It may take days to respond, compared to minutes for innate immunity. Adaptive immunity is highly specific and directed.

B cells and T cells carry out adaptive immunity. In the **humoral immune response**, B cells produce antibodies in response to activation by T cells. (“Humor” means fluid; antibodies are carried in fluids.) In the **cellular immune response**, T cells produce cytokines and activate other cells. B and T cells differentiate in the bone marrow and migrate to the lymph nodes, spleen, and thymus gland, as well as circulate in the blood and tissue fluid.

The adaptive arm of the immune system has three basic characteristics. It is *diverse*, vanquishing many types of pathogens. It is *specific*, distinguishing the cells and molecules

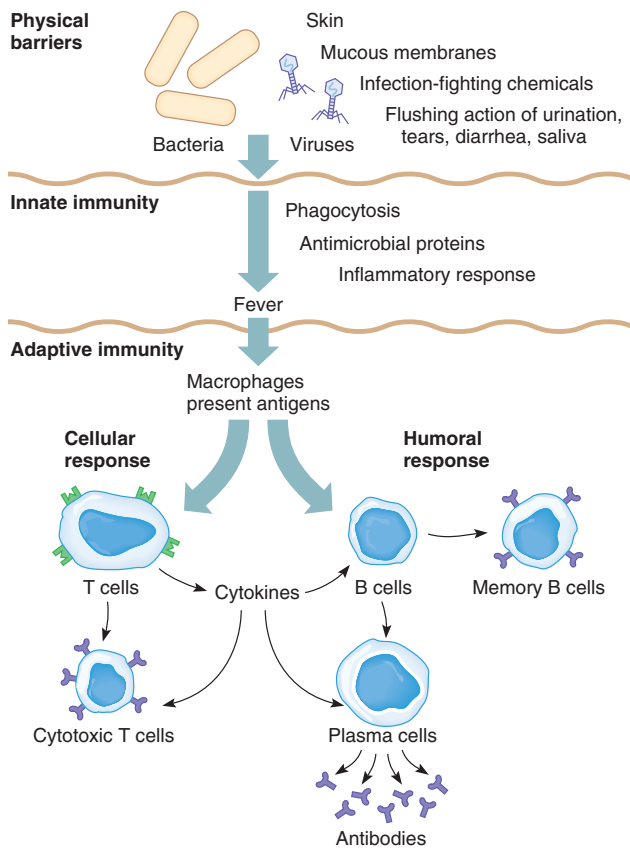


Figure 17.4 Levels of immune protection. Disease-causing organisms and viruses (pathogens) first must breach physical barriers, then nonspecific cells and molecules attack in the innate immune response. If this is ineffective, the adaptive immune response begins: Antigen-presenting cells stimulate T cells to produce cytokines, which activate B cells to divide and differentiate into plasma cells, which secrete antibodies. Once activated, these specific cells “remember” the pathogen, allowing faster responses to subsequent encounters.

that cause disease from those that are harmless. The immune system also *remembers*, responding faster to a subsequent encounter with a foreign antigen than it did the first time. The first assault initiates a **primary immune response**. The second assault, based on the system’s “memory,” is a **secondary immune response**. This is why we get some infections, such as chickenpox, only once. However, upper respiratory infections and influenza recur because the causative viruses mutate, presenting a different face to our immune systems each season.

The Humoral Immune Response—B Cells and Antibodies

An antibody response begins when an antigen-presenting macrophage activates a T cell. This cell in turn contacts a B cell that has surface receptors that can bind the type of foreign antigen the macrophage presents. The immune system has so many B cells, each with different combinations of surface antigens,

that there is almost always one or more available that corresponds to a particular foreign antigen. Turnover of these cells is high. Each day, millions of B cells perish in the lymph nodes and spleen, while millions more form in the bone marrow, each with a unique combination of surface molecules.

Once the activated T cell finds a B cell match, it releases cytokines that stimulate the B cell to divide. Soon the B cell gives rise to two types of cells (**figure 17.6**). The first, **plasma cells**, are antibody factories, each secreting 1,000 to 2,000 identical antibodies per second into the bloodstream. They live only days. These cells provide the primary immune response. Plasma cells derived from different B cells secrete different antibodies.

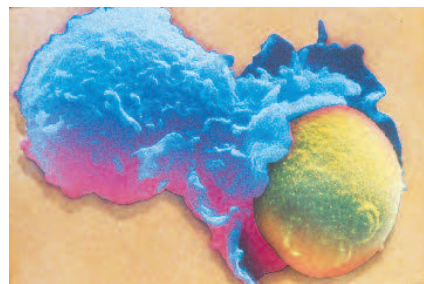


Figure 17.5 Nature's garbage collectors. A human phagocyte engulfs a yeast cell.

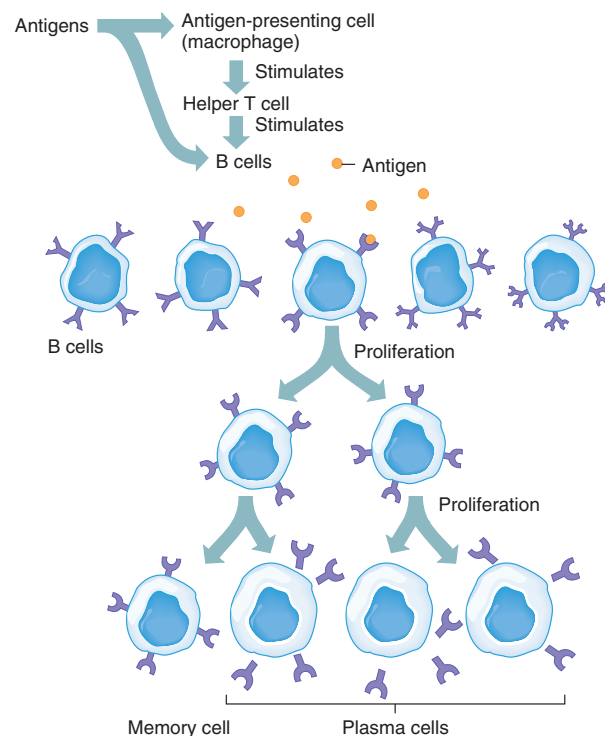


Figure 17.6 Production of antibodies. In the humoral immune response, B cells proliferate and mature into antibody-secreting plasma cells. Note that only the B cell that binds the antigen proliferates; its descendants may develop into memory cells or plasma cells. Plasma cells greatly outnumber memory cells.

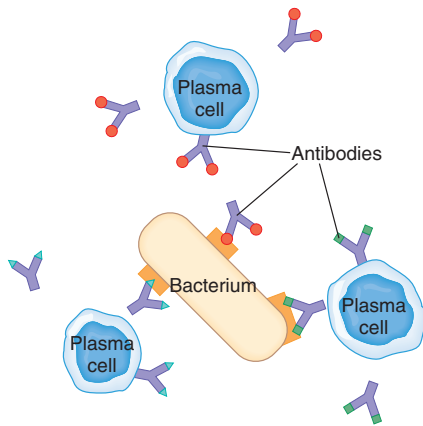


Figure 17.7 An immune response recognizes many targets. A humoral immune response is polyclonal, which means that different plasma cells produce antibody proteins that recognize and bind to different features of a foreign cell's surface.

Each type of antibody corresponds to a specific part of the pathogen, like hitting a person in different parts of the body. This multi-pronged attack is called a polyclonal antibody response (figure 17.7). The second type of B cell descendant, **memory cells**, are far fewer and usually dormant. They respond to the foreign antigen faster and with more force should it appear again. This is a secondary immune response. Memory B cells are what enabled survivors of the 1918 flu pandemic to resist infection.

An antibody molecule is built of several polypeptides and is therefore encoded by several genes. The simplest type of antibody molecule is four polypeptide chains connected by

disulfide (sulfur-sulfur) bonds, forming a shape like the letter Y (figure 17.8). A large antibody molecule might consist of three, four, or five such Ys joined.

In a Y-shaped antibody subunit, the two longer polypeptides are called **heavy chains**, and the other two **light chains**. The lower portion of each chain is an amino acid sequence that is very similar in all antibody molecules, even in different species. These areas are called constant regions, and they provide the activity of the antibody. The amino acid sequences of the upper portions of each polypeptide chain, the variable regions, can differ greatly among antibodies. These parts provide the specificities of particular antibodies to particular antigens.

Antibodies can bind certain antigens because of the three-dimensional shapes of the tips of the variable regions. These specialized ends are **antigen binding sites**, and the parts that actually contact the antigen are called **idiotypes**. The parts of the antigens that idiotypes bind are **epitopes**. An antibody contorts to form a pocket around the antigen.

Antibodies have several functions. Antibody-antigen binding may inactivate a pathogen or neutralize the toxin it produces. Antibodies can clump pathogens, making them more visible to macrophages, which then destroy them. Antibodies also activate complement, extending the innate immune response. In some situations, the antibody response can be harmful.

Antibodies are of five major types, distinguished by where they act and what they do (table 17.2). (Antibodies are also called immunoglobulins, abbreviated *Ig*.) Different antibody types predominate in different stages of an infection.

The human body can manufacture seemingly limitless varieties of antibodies, though the genome has a limited number of antibody genes. This great diversity is possible because parts of

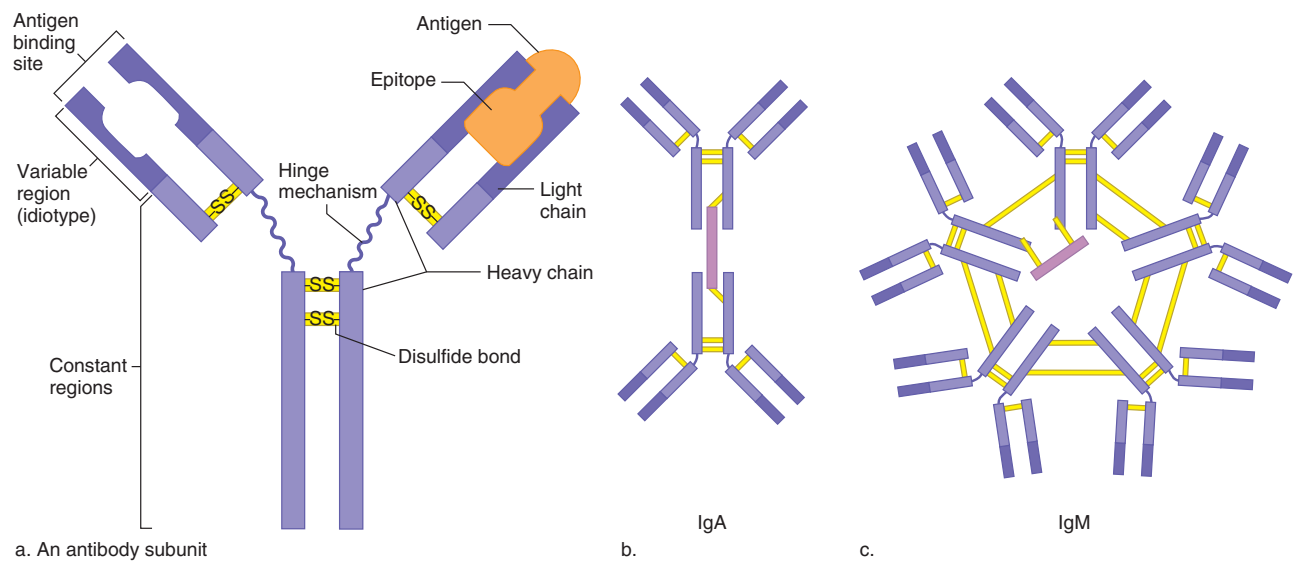


Figure 17.8 Antibody structure. The simplest antibody molecule (a) consists of four polypeptide chains, two heavy and two light, joined by pairs of sulfur atoms that form disulfide bonds. Part of each polypeptide chain has a constant sequence of amino acids, and the remainder varies. The tops of the Y-shaped molecules form antigen binding sites. (b) IgA consists of two Y-shaped subunits, and IgM (c) consists of five subunits.

Table 17.2 Types of Antibodies

Type*	Location	Functions
IgA	Milk, saliva, urine, and tears; respiratory and digestive secretions	Protects against pathogens at points of entry into body
IgD	On B cells in blood	Stimulates B cells to make other types of antibodies, particularly in infants
IgE	In secretions with IgA and in mast cells in tissues	Acts as receptor for antigens that cause mast cells to secrete allergy mediators
IgG	Blood plasma and tissue fluid; passes to fetus	Protects against bacteria, viruses, and toxins, especially in secondary immune response
IgM	Blood plasma	Fights bacteria in primary immune response; includes anti-A and anti-B antibodies of ABO blood groups

The letters A, D, E, G, and M refer to the specific conformation of heavy chains characteristic of each class of antibody.

different antibody genes combine. During the early development of B cells, sections of their antibody genes move to other chromosomal locations, creating new genetic instructions for antibodies.

The assembly of antibody molecules is like putting together many different outfits from the contents of a closet containing 200 pairs of pants, a drawer containing fifteen different shirts, and four belts. Specifically, each variable region of a heavy chain and a light chain consists of three sections, called V (for variable), D (for diversity), and J (for joining). The V, D, and J genes—several of each—for the heavy chains are on chromosome 14, and the corresponding genes for the light chains are on chromosomes 2 and 22. C (constant) genes encode the constant regions of each heavy and light chain. A promoter sequence precedes the V genes and an enhancer sequence precedes the C genes. These control sequences oversee the mixing and matching of the V, D, and J genes. **Figure 17.9** shows how the genetic instructions for the antibody parts are combined in different ways to encode the heavy and light polypeptide chains.

Enzymes cut and paste the pieces of antibody gene parts. The number of combinations is so great that virtually any antigen that a person with a healthy immune system might encounter will elicit an immune response.

The Cellular Immune Response—T Cells and Cytokines

T cells provide the cellular immune response. It is called “cellular” because the T cells themselves travel to where they act, unlike B cells, which secrete antibodies into the bloodstream. T cells descend from stem cells in the bone marrow, then travel to the thymus gland (“T” refers to thymus). As the immature T cells, called thymocytes, migrate toward the interior of the

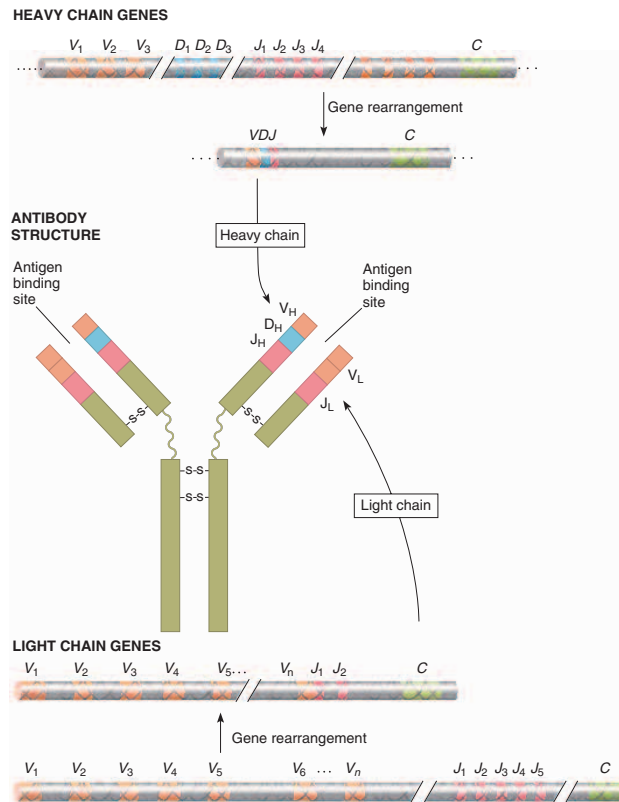


Figure 17.9 Antibody diversity. The human immune system can produce antibodies to millions of possible antigens because each polypeptide is encoded by more than one gene. That is, the many components of antibodies can combine in many ways.

thymus, they display diverse cell surface receptors. Then selection happens. As the wandering thymocytes touch lining cells in the gland that are studded with “self” antigens, thymocytes that do not attack the lining cells begin maturing into T cells, whereas those that harm the lining cells die by apoptosis—in great numbers. Gradually, T cells-to-be that recognize self persist while those that harm body cells are destroyed.

Several types of T cells are distinguished by the types and patterns of receptors on their surfaces, and by their functions. Helper T cells have many functions: they recognize foreign antigens on macrophages, stimulate B cells to produce antibodies, secrete cytokines, and activate another type of T cell called a cytotoxic T cell, (also called a killer T cell). Certain T cells may help to suppress an immune response when it is no longer required. The cytokines that helper T cells secrete include interleukins, interferons, tumor necrosis factor, and colony stimulating factors, which stimulate white blood cells in bone marrow to mature (**table 17.3**). Cytokines interact with and signal each other, sometimes in complex cascades.

Distinctive surfaces distinguish subsets of helper T cells. Certain antigens called cluster-of-differentiation antigens, or CD antigens, enable T cells to recognize foreign antigens displayed on macrophages. One such cell type, called a CD4

Cytokine	Function
Colony stimulating factors	Stimulate bone marrow to produce lymphocytes
Interferons	Block viral replication, stimulate macrophages to engulf viruses, stimulate B cells to produce antibodies, attack cancer cells
Interleukins	Control lymphocyte differentiation and growth, cause fever that accompanies bacterial infection
Tumor necrosis factor	Stops tumor growth, releases growth factors, stimulates lymphocyte differentiation, dismantles bacterial toxins

helper T cell, is an early target of HIV. Considering the critical role helper T cells play in coordinating immunity, it is little wonder that HIV infection ultimately topples the entire system, a point we will return to soon.

Cytotoxic T cells lack CD4 receptors but have CD8 receptors. These cells attack virally infected and cancerous cells by attaching to them and releasing chemicals. They do this by linking two surface peptides to form structures called T cell receptors that bind foreign antigens. When a cytotoxic T cell encounters a nonself cell—a cancer cell, for example—the T cell receptors draw the two cells into physical contact. The T cell then releases a protein called perforin, which pierces the cancer cell's plasma membrane, killing it (**figure 17.10**). Cytotoxic T cell receptors also attract cells that are covered with certain viruses, destroying the cells before the viruses on them can enter, replicate, and spread the infection. Cytotoxic T cells continually monitor body cells, recognizing and eliminating virally infected and tumor cells.

Table 17.4 summarizes types of immune system cells.

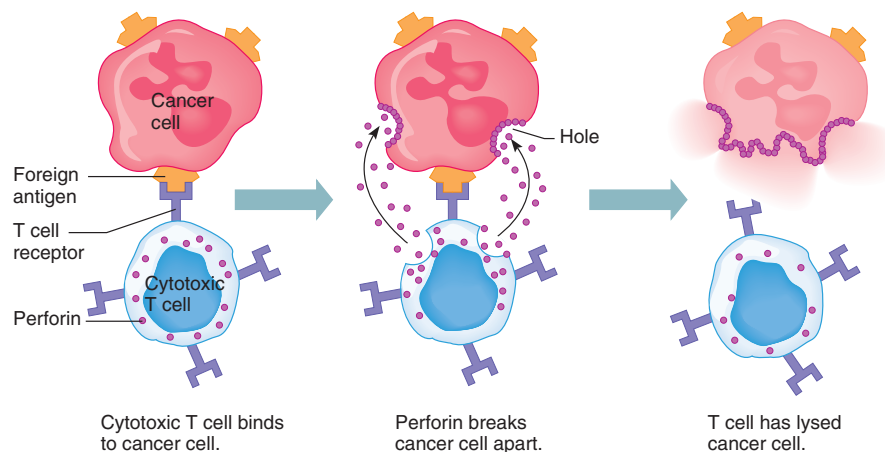


Figure 17.10 **Death of a cancer cell.** A cytotoxic T cell binds to a cancer cell and injects perforin, a protein that pierces (lyses) the cancer cell's plasma membrane. The cancer cell dies, leaving debris that macrophages clear away.

Key Concepts

1. The immune system consists of physical barriers; an innate immune response of inflammation, phagocytosis, complement, collectins, and cytokines; and an adaptive immune response that is diverse, specific, and remembers.
2. In the humoral immune response, stimulated B cells divide and differentiate into plasma cells and memory cells. A plasma cell secretes abundant antibodies of a single type. Antibodies are Y-shaped polypeptides, each with two light and two heavy chains, each with a constant and a variable region. The tips of the Y form an antigen binding site with a specific idiootype. Antibodies make foreign antigens more visible to macrophages and stimulate complement. Shuffling gene pieces generates antibody diversity.
3. In the cellular immune response, helper T cells stimulate B cells to manufacture antibodies and cytotoxic T cells to secrete cytokines. Using T cell receptors, cytotoxic T cells bind to nonself cells and virus-covered cells and burst them.

17.3 Abnormal Immunity

The immune system continually adapts to environmental change. Because the immune response is so diverse, its breakdown affects health in many ways. Immune system malfunction may be inherited or acquired, and immunity may be too weak, too strong, or misdirected. Abnormal immune responses may be multifactorial, with several genes contributing to susceptibility to infection, or caused by mutation in a single gene. Or, susceptibility to an immune disorder may reflect abnormal gene expression, as the chapter opener describes for rheumatoid arthritis.

Inherited Immune Deficiencies

The more than twenty types of inherited immune deficiencies affect innate and adaptive immunity (**table 17.5**). These conditions can arise in several ways.

In chronic granulomatous disease, neutrophils can engulf bacteria, but, due to deficiency of an enzyme called an oxidase, they cannot produce the activated oxygen compounds that kill bacteria. Because this enzyme is made of four polypeptide chains, four genes encode it, and there are four ways to inherit the disease, all X-linked. A very rare autosomal recessive form is caused by a defect in the part of the host cell that encloses bacteria.

Table 17.4 Types of Immune System Cells

Cell Type	Function
Macrophage	Presents antigens Performs phagocytosis
Mast cell	Releases histamine in inflammation Releases allergy mediators
B cell	Matures into antibody-producing plasma cell or into memory cell
T cells	
Helper	Recognizes nonself antigens presented on macrophages Stimulates B cells to produce antibodies Secretes cytokines Activates cytotoxic T cells
Cytotoxic	Attacks cancer cells and cells infected with viruses upon recognizing antigens
Natural killer	Attacks cancer cells and cells infected with viruses without recognizing antigens; activates other white blood cells
Suppressor	Inhibits antibody production

Antibiotics and gamma interferon are used to prevent bacterial infections in these patients, and the disease can be cured with a bone marrow or an umbilical cord stem cell transplant.

Mutations in genes that encode cytokines or T cell receptors impair cellular immunity, which primarily targets viruses and cancer cells. Because T cells activate the B cells that manufacture antibodies, abnormal cellular immunity (T cell function) disrupts humoral immunity (B cell function). Mutations in the genes that encode antibody segments, that control how the segments join, or that direct maturation of B cells mostly impair immunity against bacterial infection. Inherited immune deficiency can also result from defective B cells, which usually increases vulnerability to certain bacterial infections.

Severe combined immune deficiencies (SCIDs) affect both humoral and cellular immunity. About half of SCID cases are X-linked. In a less severe form, the individual lacks B cells but has some T cells. Before antibiotic drugs became available, individuals with this form of SCID died before age 10 of overwhelming bacterial infection. In a more severe form of X-linked SCID, lack of B and T cells causes death by 18 months of age, usually of severe and diverse infections. Gene therapy for SCID is discussed further in chapter 20.

A young man named David Vetter taught the world about the difficulty of life without immunity years before AIDS arrived. David had an autosomal recessive form of SCID that caused him to be born without a thymus gland. His T cells could not mature and activate B cells, leaving him defenseless in a germ-filled world. Born in Texas in 1971, David spent his short life in a vinyl bubble, awaiting a treatment that never came (figure 17.11). As he reached adolescence, David wanted to leave his bubble. A bone marrow transplant that would have enabled him to do so was unsuccessful. David left the bubble, but died within days of a massive infection.

Acquired Immune Deficiency Syndrome

AIDS is not inherited, but acquired by infection with HIV, a virus that gradually shuts down the immune system. The effect of HIV on a human body is especially astounding because the virus is so simple. Its genome is a millionth the size of ours, and its nine genes, consisting of about 9,000 RNA bases, encode only fifteen proteins! But HIV commandeers more than 200 human proteins as it invades the immune system. Infection begins as HIV enters macrophages, impairing this first line of defense. In these cells and later in helper T cells, the virus adheres with its surface protein, called gp120, to two coreceptors on the host cell surface, CD4 and CCR5 (figure 17.12). Another glycoprotein, gp41, anchors gp120 molecules into the viral envelope. When the virus binds both coreceptors,

Table 17.5 Inherited Immune Deficiencies

Disease	MIM	Inheritance*	Defect
Chronic granulomatous disease	306400	ar, AD, Xlr	Abnormal phagocytes can't kill engulfed bacteria
Immune defect due to absence of thymus	242700	ar	No thymus, no T cells
Neutrophil immunodeficiency syndrome	608203	ar	Deficiencies of T cells, B cells, and neutrophils
SCID			
Adenosine deaminase deficiency	102700	ar	No T or B cells
Adenosine deaminase deficiency with sensitivity to ionizing radiation	602450	ar	No T, B, or natural killer cells
IL-2 receptor mutation	300400	Xlr	No T, B, or natural killer cells
X-linked lymphoproliferative disease	308240	Xlr	Absence of protein that enables T cells to bind B cells

*ar = autosomal recessive

SCID = severe combined immune deficiency

AD = autosomal dominant

Xlr = X-linked recessive



Figure 17.11 David Vedder, the original “bubble boy,” was born without a thymus gland. Because his T cells could not mature, he was virtually defenseless against infection.

virus and cell surface contort in a way that enables viruses to enter the cell. Once in the cell, reverse transcriptase copies the viral RNA into DNA, which replicates to form a DNA double helix. This enters the nucleus and inserts into a chromosome. The viral DNA sequences are transcribed and translated, and the cell fills with viral pieces, which are assembled into

complete new viral particles that eventually bud from the cell (figure 17.13).

Once helper T cells start to die at a high rate, bacterial infections begin, because B cells aren’t activated to produce antibodies. Much later in infection, HIV variants arise that can bind to a receptor called CXCR4 on cytotoxic T cells, killing them. Loss of these cells renders the body very vulnerable to viral infections and cancer.

HIV replicates quickly, changes quickly, and can hide. The virus mutates easily because it cannot repair replication errors and errors happen frequently—1 per every 5,000 or so bases—because of the “sloppiness” of reverse transcriptase in copying viral RNA into DNA. The immune system cannot keep up; antibodies against one viral variant are useless against the next. For several years, the bone marrow produces 2 billion new T and B cells a day. A million to a billion new HIV particles bud daily from infected cells.

So genetically diverse is the population of HIV in a human host that, within days of the initial infection, variants arise that resist the drugs used to treat AIDS (see figure 15.12). HIV’s changeable nature is why combining drugs with different actions is the most effective way to slow the disease into a chronic, lifelong, but treatable illness, instead of a killer (table 17.6). Several classes of drugs have cut the death rate from AIDS dramatically. They work at different points of infection: blocking binding or entry of the virus into T cells, replicating viral genetic material, and processing viral proteins.

Clues to developing new drugs to treat HIV infection come from people at high risk who resist infection. Researchers identified variants of four receptors or the molecules that bind to them that block HIV from entering cells by looking at the DNA of people who had unprotected sex with many partners but who never became infected. Some of them were homozygous recessive for a 32-base deletion in the *CCR5* gene. Their *CCR5* coreceptors were too stunted to reach the cell’s surface, so HIV had nowhere to dock. Heterozygotes, with one copy of the deletion, can become infected, but they remain healthy for several years longer than people who do not have the deletion. Curiously, the same *CCR5* mutation may have enabled people to survive plague in Europe during the Middle Ages. Apparently both the virus that causes AIDS and the bacterium that causes plague use the same portal into a human cell.

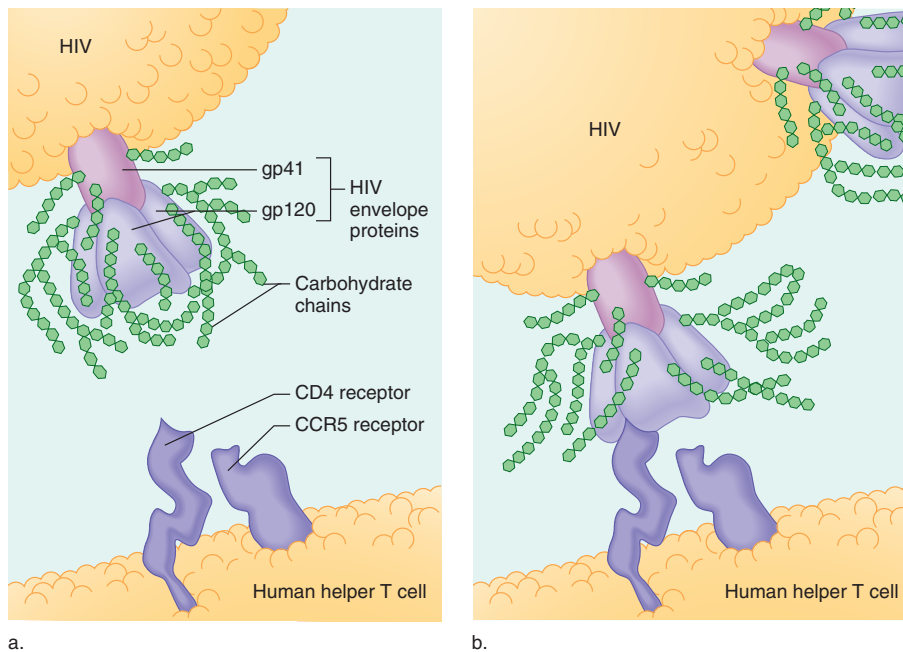


Figure 17.12 HIV binds to a helper T cell. (a) The part of HIV that binds to helper T cells is called gp120 (gp stands for glycoprotein). (b) The carbohydrate chains that shield the protein portion of gp120 move aside as they approach the cell surface, and the viral molecule can now bind to a CD4 receptor. Binding to the CCR5 receptor is also necessary. Then the viral envelope fuses with the plasma membrane and the virus enters. (The size of HIV is greatly exaggerated.)

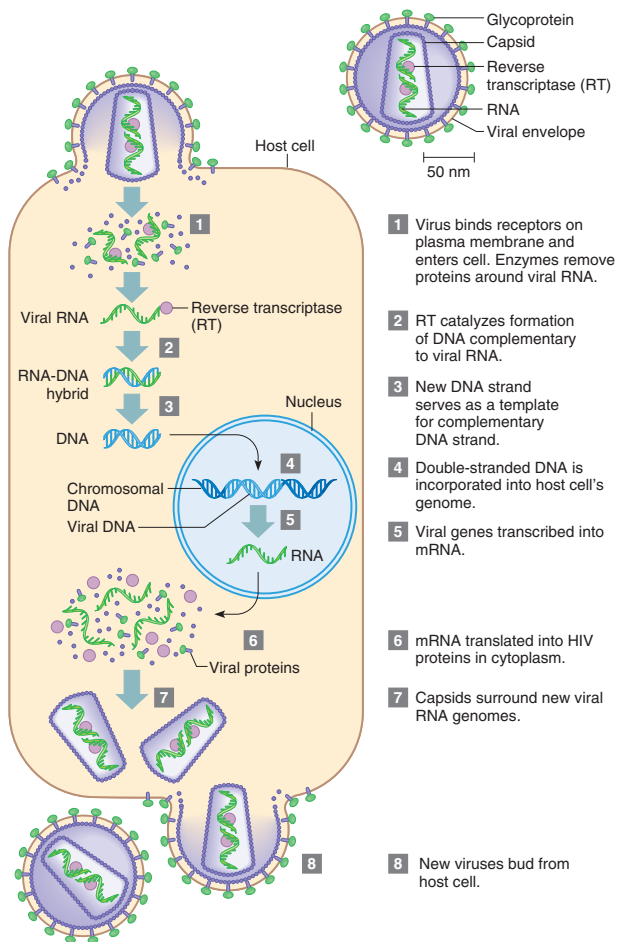


Figure 17.13 How HIV infects. HIV integrates into the host chromosome, then commandeers transcription and translation, ultimately producing more virus particles.

Autoimmunity

In **autoimmunity**, the immune system produces antibodies that attack the body's own tissues. These antibodies are called **autoantibodies**. About 5 percent of the population has an autoimmune disorder. The signs and symptoms resulting from autoimmune disorders reflect the cell types under attack (**table 17.7**).

Table 17.6 Anti-HIV Drugs

Drug Type	Mechanism
Reverse transcriptase inhibitor	Blocks copying of viral RNA into DNA
Protease inhibitor	Blocks shortening of certain viral proteins
Fusion inhibitor	Blocks ability of HIV to bind a cell
Entry inhibitor	Blocks ability of HIV to enter a cell

Most autoimmune disorders are not inherited as single-gene diseases. However, the fact that different autoimmune disorders affect members of the same family, and may respond to the same drugs, suggests that these conditions stem from shared susceptibilities. For example, in autoimmune polyendocrinopathy syndrome type I (MIM 240300), caused by a mutation in a single gene on chromosome 21, autoantibodies attack endocrine glands in a sequence, so that different members of a family may have very different symptoms. However, the genetics of autoimmunity is usually more complex than this. Genome-wide association studies have identified dozens of genes that are each associated with more than one autoimmune disorder. On the other hand, an autoimmune disorder may result from the actions of variants of several genes that each contributes to susceptibility, perhaps in the presence of a specific environmental trigger such as diet. This is the case for the digestive disorder Crohn's disease, which has been associated with thirty-two genome regions.

Some of the more common autoimmune disorders may actually arise in several ways when parts of the immune response are overactive. This is the case for systemic lupus erythematosus, better known as "lupus." The name "lupus" comes from the characteristic butterfly-shaped rash on the cheeks, but the "systemic" is actually a more important description because the disease produces autoantibodies that affect the connective tissue of many organs. These are the kidneys, joints, lungs, brain, spinal cord, and the heart and blood vessels. A person may need dialysis when the kidneys are involved, blood pressure medication to counter increasing pressure in the lungs, and drugs to minimize buildup of fatty deposits on artery walls. Lupus can also cause strokes, memory loss, fever, seizures, headache, and psychosis.

Lupus involves several aspects of the immune response, including cell surface characteristics, secretion of interferons, production of autoantibodies, activation of B and T cells, antigen presentation, adhesion of immune system cells to blood vessel linings, inflammation, removal of complexes of immune cells and foreign antigens, and cytokine production. Therefore, it isn't surprising that variants of at least ten different genes can predispose a person to developing this condition. Perhaps inheriting susceptibility in three or four of the genes causes lupus.

How does the immune system turn against itself? Autoimmunity may arise in several ways:

- A virus replicating in a cell incorporates proteins from the cell's surface onto its own. When the immune system "learns" the surface of the virus to destroy it, it also learns to attack human cells that normally bear the protein.
- Some cells that should have died in the thymus somehow escape the massive die-off, persisting to attack "self" tissue later on.
- A nonself antigen coincidentally resembles a self antigen, and the immune system attacks both. In rheumatic fever, for example, antigens on heart valve cells resemble those on *Streptococcus* bacteria; antibodies produced to fight a strep throat also attack the heart valve cells.
- If X inactivation is skewed, a female may have a few cells that express the X chromosome genes of one parent. The immune system may respond to these cells as foreign

Table 17.7 Autoimmune Disorders

Disorder	Symptoms	Autoantibodies against
Diabetes mellitus (type 1)	Thirst, hunger, weakness, weight loss	Pancreatic beta cells
Graves disease	Restlessness, weight loss, irritability, increased heart rate and blood pressure	Thyroid gland cells
Hemolytic anemia	Fatigue, weakness	Red blood cells
Multiple sclerosis	Weakness, poor coordination, failing vision, disturbed speech	Myelin in the white matter of the central nervous system
Myasthenia gravis	Muscle weakness	Neurotransmitter receptors on skeletal muscle cells
Rheumatic fever	Weakness, shortness of breath	Heart valve cells
Rheumatoid arthritis	Joint pain and deformity	Cells lining joints
Systemic lupus erythematosus	Red facial rash, fever, weakness, joint pain	Connective tissue
Ulcerative colitis	Lower abdominal pain	Colon cells

if they have surface antigens that are not also on the majority of cells. Skewed X inactivation may explain why some autoimmune disorders are much more common in females.

Reading 17.2 highlights a special situation in which two immune systems must co-exist—pregnancy.

Allergies

An allergy is an immune system response to a substance, called an allergen, that does not actually present a threat. Many allergens are particles small enough to be carried in the air and enter a person's respiratory tract. The size of the allergen may determine the type of allergy. For example, grass pollen is large and remains in the upper respiratory tract, where it causes hayfever. But allergens from house dust mites, cat dander, and cockroaches are small enough to infiltrate the lungs, triggering asthma. Asthma is a chronic disease in which contractions of the airways, inflammation, and accumulation of mucus block air flow.

Both humoral and cellular immunity take part in an allergic response (**figure 17.14**). Antibodies of class IgE bind to mast cells, sending signals that cause the mast cells to open and release allergy mediators such as histamine and heparin. Allergy mediators cause inflammation, with symptoms that may include runny eyes from hay fever, narrowed airways from asthma, rashes, or the overwhelming body-wide allergic reaction called anaphylactic shock. Allergens also activate a class of helper T cells that produce a particular mix of cytokines whose genes are clustered

on chromosome 5q. Regions of chromosomes 12q and 17q have genes that control IgE production.

The fact that allergies have become very common only during the past century suggests a much stronger environmental than genetic component. Still, people inherit susceptibilities to allergy. Twin studies of various allergies reveal about a 75 percent concordance, and isolated populations with a great deal of inbreeding tend to have a high prevalence of certain allergies.

The allergies that people suffer today may be a holdover of an immune function that was important in the past. Evidence for this idea is that people with allergies have higher levels of white blood cells called eosinophils than do others, and these cells fight parasitic infections that are no longer common. In a more general sense, because allergies are more prevalent in developed

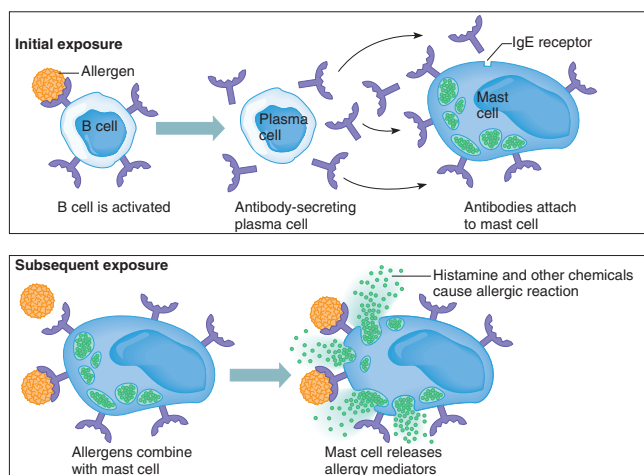


Figure 17.14 Allergy. In an allergic reaction, an allergen such as pollen activates B cells, which divide and give rise to antibody-secreting plasma cells. The antibodies attach to mast cells. When the person encounters allergens again, the allergens combine with the antibodies on the mast cells, which then burst, releasing the chemicals that cause itchy eyes and a runny nose.

Reading 17.2

A Special Immunological Relationship: Mother-to-Be and Fetus

The immune system recognizes “self” cell surfaces and protects the body from foreign, “nonself” cells and molecules. This is very helpful when the nonself triggers are parts of infecting bacteria, but what tempers the immune system of a pregnant woman to accept cells from her fetus? Half of a fetal genome comes from the father, and so fetal cell surfaces likely include some antigens from him that would be “foreign” to the mother-to-be. Similarly, some of her antigens might be foreign to the fetus. Yet, pregnant woman and fetus routinely swap cells.

The immune system must have evolved a tolerance between woman and fetus, or else we would not be here! We do not understand this phenomenon well, but following are three examples of the immunological “crosstalk” between a pregnant woman and her fetus.

“T Regs”

Samples of lymph nodes from fetuses mid-pregnancy indicate that up to 1 percent of the cells are maternal. The woman’s cells stimulate the fetal immune system to produce “regulatory T cells,” called “T Regs,” which dampen the immune response. The maternal immune system does the same. In one experiment, fetal lymph node samples did not react against cells from the mother unless the regulatory T cells were removed. Children retain these cells for several years. It may be possible to stimulate production of T Regs later in life to help a recipient’s body accept an organ transplant.



Figure 1 An autoimmune disorder—**maybe**. Scleroderma hardens the skin. Some cases appear to be caused by a long-delayed immune response to cells retained from a fetus decades earlier.

Scleroderma

People who have scleroderma describe the condition as “the body turning to stone.” The skin hardens, causing a mask-like appearance, and the skin elsewhere looks more like armor (**figure 1**). Scleroderma usually begins in middle age, and affects mostly women. It was long thought to be an autoimmune disorder, but the discovery of Y chromosomes in skin cells from scleroderma patients who are mothers of sons revealed a very different source of the illness—lingering cells from a fetus. Cells from female fetuses can presumably have the same effect but cannot be distinguished from the mother on the basis of a sex chromosome check.

The degree of genetic difference between a mother and a son may play a role in development of scleroderma. Mothers who have the condition tend to have cell surfaces that are more similar to those of their sons than mothers who do not have scleroderma. Perhaps the similarity of cell surfaces enabled the fetal cells to escape destruction by the mother’s immune system.

Rh Incompatibility

“Rh,” the rhesus factor discovered in rhesus monkeys, is a blood group (MIM 111700). A person is Rh⁺ if red blood cells have a surface molecule called the RhD antigen. Rh type is important when an Rh⁺ man and an Rh⁻ woman conceive a child who is Rh⁺ (**figure 2**). The woman’s immune system manufactures antibodies against the few fetal cells that enter her bloodstream. Not enough antibodies form to harm the fetus that sets off the reaction. However, the number of antibodies continues to increase. If she carries another Rh⁺ fetus, the antibodies can attack the fetal blood supply, causing a potentially fatal condition called hemolytic disease of the fetus and newborn. It can be treated at birth with a transfusion of Rh⁻ blood.

Fortunately, natural and medical protections make this complication rare today. If a woman’s ABO blood type is O and the fetus is A or B, her anti-A or anti-B antibodies attack the fetal cells in her circulation before her immune system produces anti-Rh antibodies. Also, if a pregnant woman alerts her physician to a potential incompatibility, she can be given a substance called RhoGAM, which is antibody against the Rh antigen that shields fetal cells so her system does not manufacture the harmful antibodies. When she becomes pregnant again, fetal DNA in her circulation can be tested to see if it is Rh⁻ or Rh⁺. If the second fetus is Rh⁻, she does not need RhoGAM.

A first Rh⁺ fetus developing in an Rh⁻ mother can be affected if her blood has been exposed to Rh⁺ cells in any of several circumstances: amniocentesis, a blood transfusion, an ectopic (tubal) pregnancy, a miscarriage, or an abortion.

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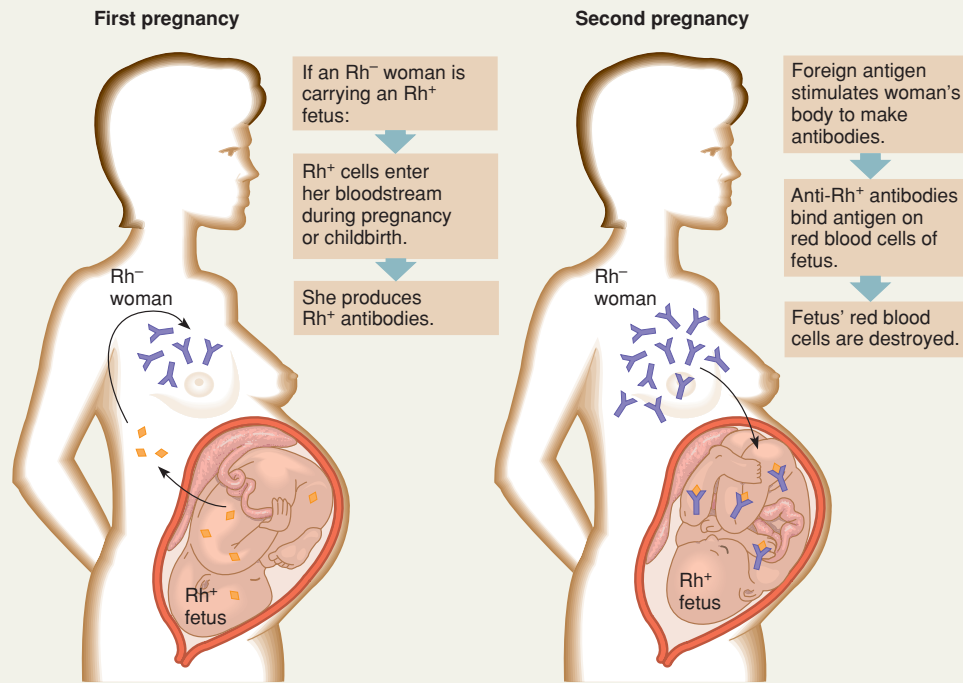


Figure 2 Rh incompatibility. Fetal cells entering the pregnant woman's bloodstream can stimulate her immune system to make anti-Rh antibodies, if the fetus is Rh⁺ and she is Rh⁻. A drug called RhoGAM prevents attacks on subsequent fetuses.

nations and have become more common since the introduction of antibiotic drugs, allergies may result from a childhood relatively free of infection, compared to times past—almost as if the immune system is reacting to being underutilized. This idea that allergies stem from an environment too clean to have stimulated the immune system very much is called the hygiene hypothesis.

Key Concepts

1. Inherited immune deficiencies affect innate and adaptive immunity.
2. HIV is a retrovirus that injects its RNA into host cells by binding coreceptors. Reverse transcriptase then copies viral RNA into DNA. HIV uses the cell's protein synthesis machinery to mass produce itself; then the cell releases virus. HIV continually mutates, becoming resistant to drugs. HIV replicates very rapidly, and T cell production matches it until the immune response is overwhelmed.
3. In autoimmune disorders, autoantibodies attack healthy tissue.
4. An allergic reaction results from a misdirected immune response. Allergens bind to IgE antibodies on mast cells, which release allergy mediators.

17.4 Altering Immunity

Medical technology can alter or augment immune system functions in various ways. Vaccines trick the immune system into acting early. Antibiotic drugs, which are substances derived from organisms such as fungi and soil bacteria, have been used for decades to assist an immune response. Cytokines and altered antibodies are used as drugs to treat a variety of conditions. Transplants require suppression of the immune system so that the body will accept a nonself replacement body part.

Vaccines

A **vaccine** is an inactive or partial form of a pathogen that stimulates the immune system to alert B cells to produce antibodies. When the person then encounters the natural pathogen, a secondary immune response ensues, even before symptoms arise. Vaccines consisting of entire viruses or bacteria can, rarely, cause illness if they mutate to a pathogenic form. This was a risk of the smallpox vaccine. A safer vaccine uses only the part of the pathogen's surface that elicits an immune response. Vaccines against different illnesses can be combined into one injection, or the genes encoding antigens from several pathogens can be inserted into a harmless virus and delivered as a "super vaccine."

Vaccine technology dates back to the eleventh century in China. Because people saw that those who recovered from smallpox never got it again, they crushed scabs from pox into a powder that they inhaled or rubbed into pricked skin. In 1796, the wife of a British ambassador to Turkey witnessed the Chinese method of vaccination and mentioned it to English country physician Edward Jenner. Intrigued, Jenner was vaccinated the Chinese way, and then thought of a different approach.

It was widely known that people who milked cows contracted a mild illness called cowpox, but did not get smallpox. The cows became ill from infected horses. Since the virus seemed to jump from one species to another, Jenner wondered whether exposing a healthy person to cowpox lesions might protect against smallpox. A slightly different virus causes cowpox, but Jenner's approach worked, leading to development of the first vaccine (the word comes from the Latin *vacca*, for "cow").

Jenner tried his first vaccine on a volunteer, 8-year-old James Phipps. Jenner dipped a needle in pus oozing from a small cowpox sore on a milkmaid named Sarah Nelmes, then scratched the boy's arm with it. He then exposed the boy to people with smallpox. Young James never became ill. Eventually, improved versions of Jenner's smallpox vaccine eradicated a disease that once killed millions (**figure 17.15**). By the 1970s, vaccination became unnecessary. However, several nations have resumed smallpox vaccination, as section 17.5 discusses.

Most vaccines are injections. New delivery methods include nasal sprays (flu vaccine) and genetically modified fruits and vegetables. A banana as a vaccine makes sense in theory,



Figure 17.15 Smallpox: Gone? This boy is one of the last victims of smallpox, which has not naturally infected a human since 1977. Because many doctors are unfamiliar with smallpox, and people are no longer vaccinated, an outbreak would be a major health disaster.

but in practice it is difficult to obtain a uniform product. Edible plants are grown from cells that are given genes from pathogens that encode the antigens that evoke an immune response. When the plant vaccine is eaten, the foreign antigens stimulate phagocytes beneath the small intestinal lining to “present” the antigens to nearby T cells. From here, the antigens go to the bloodstream, where they stimulate B cells to divide to yield plasma cells that produce IgA. These antibodies coat the small intestinal lining, protecting against pathogens in food. Current research focuses on converting plant-based vaccines into powders so that doses can be regulated—but this counters the original goal of feeding bananas to easily immunize babies.

Whatever the form of vaccine, it is important that a substantial proportion of a population be vaccinated to control an infectious disease. This establishes “herd immunity”—that is, if unvaccinated people are rare, then if the pathogen appears, it does not spread, because so many people are protected. If the population includes unvaccinated individuals who come into contact, the disease can spread.

Immunotherapy

Immunotherapy amplifies or redirects the immune response. It originated in the nineteenth century to treat disease. Today, a few immunotherapies are in use, with more in clinical trials.

Monoclonal Antibodies Boost Humoral Immunity

When a B cell recognizes a single foreign antigen, it manufactures a single, or monoclonal, type of antibody. A large amount of a single antibody type could target a particular pathogen or cancer cell because of the antibody's great specificity.

In 1975, British researchers Cesar Milstein and George Köhler devised monoclonal antibody (MAb) technology, which mass-produces a single B cell, preserving its specificity and amplifying its antibody type. First, they injected a mouse with a sheep's red blood cells (**figure 17.16**). They then isolated a single B cell from the mouse's spleen and fused it with a cancerous white blood cell from a mouse. The fused cell, called a hybridoma, had a valuable pair of talents. Like the B cell, it produced large numbers of a single antibody type. Like the cancer cell, it divided continuously.

Today MAbs are made to more closely resemble natural human antibodies because the original mouse preparations caused allergic reactions. MAbs are used in basic research, veterinary and human health care, agriculture, forestry, and forensics. They can diagnose everything from strep throat to turf grass disease. In a home pregnancy test, a woman places drops of her urine onto a paper strip containing a MAb that binds hCG, the “pregnancy” hormone. The color changes if the MAb binds its target. In cancer diagnosis, if a MAb attached to a fluorescent dye and injected into a patient or applied to a sample of tissue or body fluid binds its target—an antigen found mostly or only on cancer cells—fluorescence indicates disease. MAbs linked to radioactive isotopes or to drugs deliver treatment to cancer cells. The MAb drug Herceptin blocks receptors on certain breast cancer cell surfaces, preventing them from receiving signals to divide.

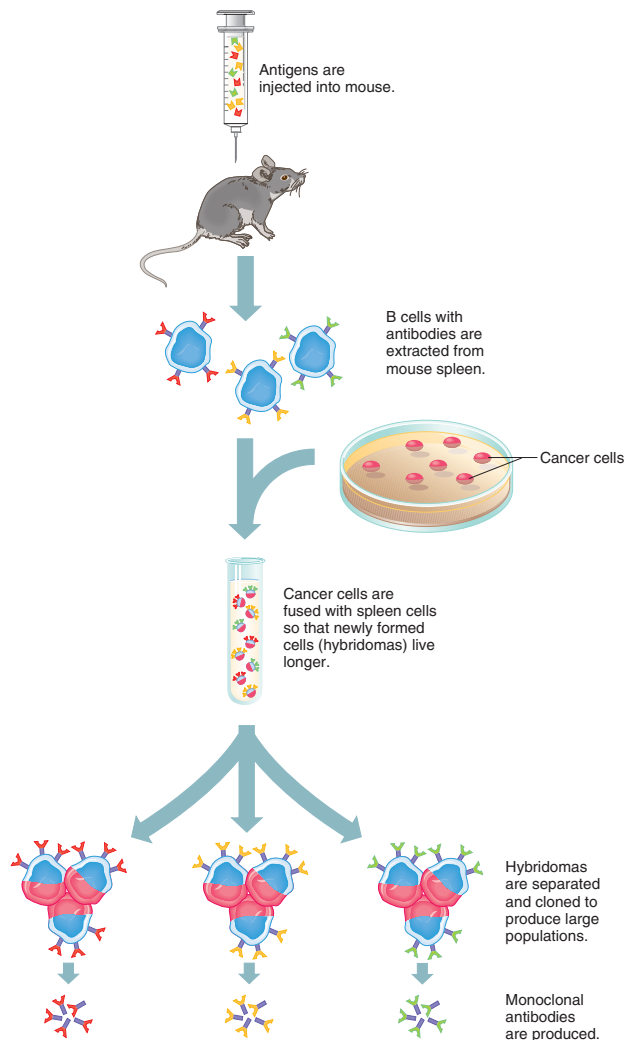


Figure 17.16 Monoclonal antibody technology. Monoclonal antibodies are pure preparations of a single antibody type that recognize a single antigen type. They are useful in diagnosing and treating disease because of their specificity.

Cytokines Boost Cellular Immunity

As coordinators of immunity, cytokines are used to treat a variety of conditions. However, it has been difficult to develop these body chemicals into drugs because they remain active only for short periods. They must be delivered precisely where they are needed, or overdose or side effects can occur.

Interferon (IF) was the first cytokine tested on a large scale. Various interferons are used to treat cancer, genital warts, and multiple sclerosis. Interleukin-2 (IL-2) is a cytokine that is administered intravenously to treat kidney cancer recurrence. Colony stimulating factors, which cause immature white blood cells to mature and differentiate, are used to boost white blood cell levels in people with suppressed immune systems, such as individuals with AIDS or those receiving cancer chemotherapy. Treatment with these

factors enables a patient to withstand higher doses of a conventional drug.

Because excess of another cytokine, tumor necrosis factor (TNF), underlies some disorders, blocking its activity treats some conditions. The drug Enbrel, for example, consists of part of a receptor for TNF. Taking it prevents TNF from binding to cells that line joints, relieving arthritis. Excess TNF in rheumatoid arthritis prevents the joint lining cells from secreting lubricating fluid.

Transplants

When a car breaks down, replacing the damaged part often fixes the trouble. The same is sometimes true for the human body. Hearts, kidneys, livers, lungs, corneas, pancreases, skin, and bone marrow are routinely transplanted, sometimes several organs at a time. Although transplant medicine had a shaky start, many problems have been solved. Today, thousands of transplants are performed annually and recipients gain years of life. The challenge to successful transplantation lies in genetics because individual inherited differences in cell surfaces determine whether the body will accept tissue from a particular donor.

Transplant Types

Transplants are classified by the relationship of donor to recipient (**figure 17.17**):

1. An autograft transfers tissue from one part of a person's body to another. A skin graft taken from the thigh to replace burned skin on the chest, or a leg vein that replaces a coronary artery, are autografts. The immune system does not reject the graft because the tissue is self. (Technically, an autograft is not a transplant because it involves only one person.)
2. An isograft is tissue from a monozygotic twin. Because the twins are genetically identical, the recipient's immune system does not reject the transplant. Ovary isografts have been performed.
3. An allograft comes from an individual who is not genetically identical to the recipient, but is a member of the same species. A kidney transplant from an unrelated donor is an allograft.
4. A xenograft transplants tissue from one species to another. (See the Bioethics Box on page 350.)

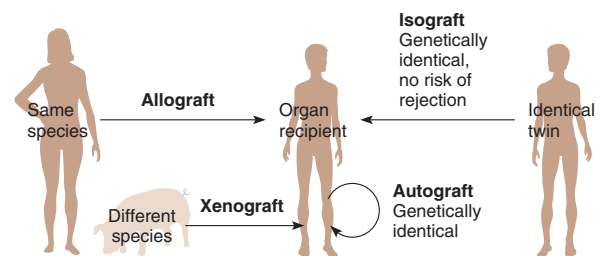


Figure 17.17 Transplant types. An autograft is within an individual. An isograft is between identical twins. An allograft is between members of the same species, and a xenograft is between members of different species.

Bioethics: Choices for the Future

Pig Parts

In 1902, a German medical journal reported an astonishing experiment. A physician, Emmerich Ullman, had attached the blood vessels of a patient dying of kidney failure to a pig's kidney set up by her bedside. The patient's immune system rejected the attachment almost immediately.

Nearly a century later, in 1997, a similar experiment took place. Robert Pennington, a 19-year-old suffering from acute liver failure and desperately needing a transplant, survived for six and a half hours with his blood circulating outside of his body through a living liver removed from a 15-week-old, 118-pound pig named Sweetie Pie. The pig liver served as a bridge until a human liver became available. But Sweetie Pie was no ordinary pig. She had been genetically modified and bred so that her cells displayed a human protein that controlled rejection of tissue transplanted from an animal of another species. Because of this slight but key bit of added humanity, plus immunosuppressant drugs, Pennington's body tolerated the pig liver's help for the few crucial hours. Baboons have also been used as sources of transplant organs (see figure).

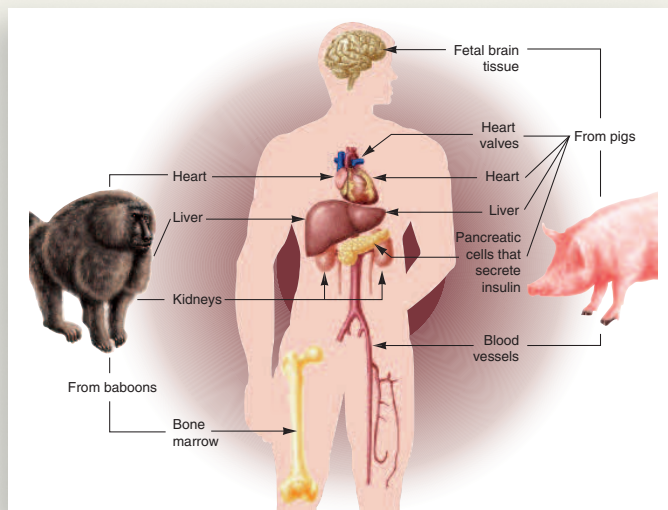
Successful xenotransplants would help alleviate the organ shortage. However, some people object to raising animals to use their organs as transplants because it requires killing the donors. One researcher counters such protests by comparing the use of animal organs as transplants to eating them.

A possible danger of xenotransplants is that people may acquire viruses from the donor organs. Viruses can "jump" species, and the outcome in the new host is unpredictable. For example, a virus called PERV—for "porcine endogenous retrovirus"—can infect human cells in culture. However, several dozen patients who received implants of pig tissue did not show evidence of PERV years later. That study, though, looked only at blood. We still do not know what effect pig viruses can have on a human body. Because many viral infections take years to cause symptoms, introducing a new infectious disease in the future could be the trade-off for using xenotransplants to solve the current organ shortage.

Questions for Discussion

- Pig parts as transplants may become necessary due to the shortage of human organs. Discuss the pros and cons of the following systems for rationing human organs:
 - First come, first serve
 - Closeness of match of cell surface antigens
 - Ability to pay
 - The importance of the recipient
 - The youngest
 - The most severely ill, who will soon die without the transplant
 - The least severely ill, who are strong enough to survive a transplant
 - Those who are not responsible for their condition, such as a nonsmoker with hereditary emphysema versus a person who has emphysema caused by smoking
- In the novel *Never Let Me Go*, Kazuo Ishiguro tells of a society in which certain people are designated as organ donors. They know that at a certain age, their organs will be removed, one by one, until they die, to provide transplants for wealthy recipients. The film *The Island* has a similar plot, except that the donors do not know their fate. In Robin Cook's novel *Chromosome Six*, a geneticist places *HLA* genes into fertilized ova from bonobos (pygmy chimps) and the animals are raised to provide organs for wealthy humans.

Read either of these books or see the film, or choose another with a transplant plot, and discuss the source of the transplants from the points of view of the donor, the recipient, the families of both, and the government.
- Discuss the issues that people might find disturbing about creating animals such as pigs or primates that have certain human molecules that make their organs more likely to be accepted as a transplant.



Baboons and pigs can provide tissues and organs for transplant.

Rejection Reactions—Or Acceptance

The immune system recognizes most donor tissue as nonself. Then, in a tissue rejection reaction, T cells, antibodies, and activated complement destroy the foreign tissue. The greater the difference between recipient and donor cell surfaces, the more rapid and severe the rejection reaction. An extreme example is the hyperacute rejection reaction against tissue transplanted from another species—the donor tissue is usually destroyed in minutes as blood vessels blacken and cut off the blood supply.

Physicians use several approaches to dampen rejection so that a transplant recipient can survive. These include closely matching the HLA types of donor and recipient and stripping donor tissue of antigens. Immunosuppressant drugs inhibit production of the antibodies and T cells that specifically attack transplanted tissue, while sparing other components of the immune system. If recipients get bone marrow stem cells from the organ donors, they need immunosuppressant drugs for only a short time. Somehow the bone marrow stem cells help the recipient's body accept the transplanted tissue. Gene expression microarrays can be used to better match donors to recipients.

Rejection is not the only problem that can arise from an organ transplant. Graft-versus-host disease develops sometimes when bone marrow transplants are used to correct certain blood deficiencies and cancers. The transplanted bone marrow, which is actually part of the donor's immune system, attacks the recipient—its new body—as foreign. Symptoms include rash, abdominal pain, nausea, vomiting, hair loss, and jaundice.

Sometimes a problem arises if a bone marrow transplant to treat cancer is too closely matched to the recipient. If the cancer returns with the same cell surfaces as it had earlier, the patient's new bone marrow is so similar to the old marrow that it is equally unable to fight the cancer. The best tissue for transplant may be different enough to control the cancer, but not so different that rejection occurs.

Key Concepts

1. Vaccines are disabled pathogens or their parts that elicit an immune response against infection by the active pathogen.
2. Immunotherapy uses immune system components to fight disease. B cells fused with cancer cells produce MAbs that target specific antigens. Cytokines boost immune function and destroy cancer cells.
3. Autografts transfer tissue from one part of a person's body to another; isografts are between identical twins; allografts are between members of the same species; and xenografts are cross-species transplants.
4. Allografts can cause tissue rejection reactions, and xenografts can set off hyperacute rejection. In graft-versus-host disease, transplanted bone marrow rejects the recipient's tissues.

17.5 A Genomic View of Immunity—The Pathogen's Perspective

Immunity against infectious disease involves interactions of two genomes—ours and the pathogen's. Human genome information is revealing how the immune system halts infectious disease. Information from pathogen genomes reveals how they make us sick.

Knowing the DNA sequence of a pathogen's genome, or the sequences of key genes, can reveal exactly how that organism causes illness in humans. This information can suggest new treatment strategies. The sequence for *Streptococcus pneumoniae*, for example, revealed instructions for a huge protein that enables the bacterium to adhere to human cells. Potential drugs could dismantle this adhesion protein.

Pathogen genome information is also used to protect against infection in an approach called reverse vaccinology. Instead of culturing hard-to-grow pathogens in the laboratory, researchers identify and use parts of genomes that encode antigens that provoke the human immune system. Researchers rapidly developed vaccines against severe acquired respiratory syndrome (SARS) in 2003, H5N1 avian flu in 2005, and H1N1 swine flu in 2009 using this strategy.

Crowd Diseases

History provides clues to the complex and ever-changing relationships between humans and our pathogens. Because adaptive immunity responds to an environmental stimulus, epidemics often followed the introduction of a pathogen into a population that had not encountered it before.

When Europeans first explored the New World, they inadvertently brought bacteria and viruses to which their immune systems had adapted. The immune systems of Native Americans, however, had never encountered these pathogens. Many people died. Smallpox decimated the Aztec population in Mexico from 20 million in 1519, when conquistador Hernán Cortés arrived from Spain, to 10 million by 1521, when Cortés returned. By 1618, the Aztec nation population had fallen to 1.6 million. The Incas in Peru and northern populations were also dying of smallpox. When explorers visited what is now the southeast United States, they found abandoned towns where natives had died from smallpox, measles, pertussis, typhus, and influenza.

The diseases that so easily killed Native Americans are known as "crowd" diseases, because they arose with the spread of agriculture and urbanization and affect many people. Crowd diseases swept Europe and Asia as expanding trade routes spread bacteria and viruses along with silk and spices. Today air travel spreads crowd diseases.

Crowd diseases tend to pass from conquerors who live in large, intercommunicating societies to smaller, more isolated and susceptible populations, not vice versa. When Columbus arrived in the New World, the large populations of Europe and Asia had existed far longer than American settlements. In

Europe and Asia, infectious diseases had time to become established and for human populations to adapt to them. In contrast, an unfamiliar infectious disease can quickly wipe out an isolated tribe, leaving no one to give the illness to new invaders.

Most crowd diseases vanish quickly, for several reasons: vaccines or treatments may stop transmission; people may alter their behaviors to avoid contracting the infection; or the disease may kill before individuals can pass it on. Sometimes, we don't know why a disease vanishes or becomes milder.

We may be able to treat and control newly evolving infectious diseases one at a time, with new drugs and vaccines. But the mutation process that continually spawns new genetic variants in microbe populations—resulting in evolution—means that new infectious diseases will continue to arise, and old ones to return or ravage new populations.

Bioweapons

It may seem incomprehensible that anyone would ever use pathogens to intentionally harm people, but it is a sad fact of history—and the present—that such bioweapons exist. Biological weapons have been around since medieval warriors catapulted plague-ridden corpses over city walls to kill the inhabitants. During the French and Indian War, the British gave Native Americans blankets intentionally contaminated with secretions from smallpox victims. Although international law banned “germ warfare” in 1925, from 1932 until 1942 Japan field-tested bacterial bioweapons in rural China, killing thousands.

In 1973, the Soviet Union established an organization called Biopreparat. Thousands of workers in 50 facilities prepared anthrax bombs and other bioweapons under the guise of manufacturing legitimate drugs, vaccines, and veterinary products. Soviet bioweapons were even more lethal than their natural counterparts. Plague bacteria, for example, were genetically modified to resist sixteen antibiotics and to manufacture a protein that strips nerve cells of their fatty coats, adding paralysis to the list of natural symptoms.

Anthrax has continued to leave its mark on the history of bioweaponry. In 1979, an accident occurred in a Soviet city then called Sverdlovsk. At Military Compound Number 19, a miscommunication among shift workers in charge of changing safety air filters caused the release of a cloud of dried anthrax spores over the city. Within weeks, more than 100 people died of anthrax. They were mostly young, healthy men who were outside on that Friday night and breathed in enough anthrax spores to cause the respiratory collapse of the inhaled form of

the illness. The government proclaimed the deaths due to eating infected meat. Workers sprayed water everywhere, reaerosolizing the spores and causing more infections. Inhalation anthrax was apparently used as a bioweapon in the United States in 2001, as the introduction to chapter 12 discusses.

Genetics underlies inhalation anthrax because the toxin the bacteria produce consists of three proteins. One protein forms a barrel-like structure that binds to macrophages and admits the other two proteins. One of these components overloads signal transduction and impairs the cell's ability to engulf pathogens. The other toxin component breaks open macrophages, which release tumor necrosis factor and interleukins.

In the United States, a small-scale bioweapons effort began in 1942. A facility at Fort Detrick in Frederick, Maryland, stored 5,000 bombs loaded with anthrax spores; a production facility for the bombs was located in Terre Haute, Indiana; and Mississippi and Utah had test sites. President Richard Nixon halted the program in 1969; he thought that conventional and nuclear weapons were a sufficient deterrent and defense. In 1972, political leaders in London, Moscow, and Washington signed the Biological Weapons Convention, an effort to prevent bioterrorism. Its protocols are being strengthened today. In September 1992, then-president Boris Yeltsin officially halted bioweapon research in the former Soviet Union.

Today bioweapons are again a threat. Bioterrorists not only know how to grow and dry pathogens, but how to control particle size to ease infection. In addition, genetic modification can alter the characteristics of a virus or bacterium intended for use as a weapon, making it even deadlier, or targeting specific types of victims.

Key Concepts

1. Knowing the genome sequence of a pathogen can reveal how it evades the human immune system.
2. Crowd diseases happen when infectious agents are introduced into a population that hasn't encountered them before.
3. Bioterrorism is the use of pathogens—either in their natural state or genetically manipulated—to kill people.

Summary

17.1 The Importance of Cell Surfaces

1. The cells and biochemicals of the immune system distinguish self from nonself, protecting the body against infections and cancer.
2. Genes encode immune system proteins, and may confer susceptibilities to certain infectious diseases.
3. An **antigen** is a molecule that elicits an immune response. Patterns of cell surface protein and glycoprotein antigens

determine blood types. **HLA** genes encode cell surface antigens that present foreign antigens to the immune system.

17.2 The Human Immune System

4. If a pathogen breaches physical barriers, the **innate immune response** produces the redness and swelling of inflammation, plus **complement**, **collectins**, and **cytokines**. The response is broad and general.

- The **adaptive immune response** is slower, specific, and has memory.
- The **humoral immune response** begins when macrophages display foreign antigens near HLA antigens. This activates **T cells**, which activate **B cells**. The B cells, in turn, give rise to plasma cells and secrete specific **antibodies**. Some B cells give rise to **memory cells**.
- An antibody is Y-shaped and has four polypeptide chains, two heavy and two light. Each antibody molecule has regions of constant amino acid sequence and regions of variable sequence.
- The tips of the Y of each subunit form antigen binding sites, which include the more specific idiotypes that bind foreign antigens at their epitopes.
- Antibodies bind antigens to form immune complexes large enough for other immune system components to detect and destroy. Antibody genes are rearranged during early B cell development, providing instructions to produce a great variety of antibodies.
- T cells carry out the **cellular immune response**. Their precursors are selected in the thymus to recognize self. Helper T cells secrete cytokines that activate other T cells and B cells. A helper T cell's CD4 antigen binds macrophages that present foreign antigens. Cytotoxic T cells release biochemicals that kill bacteria and destroy cells covered with viruses.

17.3 Abnormal Immunity

- Mutations in antibody or cytokine genes, or in genes encoding T cell receptors, cause inherited immune deficiencies. Severe combined immune deficiencies affect both branches of the immune system.
- HIV binds to the coreceptors CD4 and CCR5 on macrophages and helper T cells, and, later in infection, triggers apoptosis of cytotoxic T cells. As HIV replicates, it mutates, evading

immune attack. Falling CD4 helper T cell numbers allow opportunistic infections and cancers to flourish. People who cannot produce a complete CCR5 protein resist HIV infection.

- In an **autoimmune disease**, the body manufactures **autoantibodies** against its own cells.
- In susceptible individuals, allergens stimulate IgE antibodies to bind to mast cells, which causes the cells to release allergy mediators. Certain helper T cells release selected cytokines. Allergies may be a holdover of past immune function.

17.4 Altering Immunity

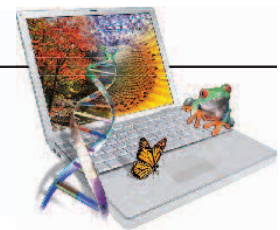
- A **vaccine** presents a disabled pathogen, or part of one, to elicit a primary immune response.
- Immunotherapy enhances or redirects immune function. Monoclonal antibodies are useful in diagnosing and treating some diseases because of their abundance and specificity. Cytokines are used to treat various conditions.
- Transplant types include autografts (within oneself), isografts (between identical twins), allografts (within a species), and xenografts (between species). A tissue rejection reaction occurs if donor tissue is too unlike recipient tissue.

17.5 A Genomic View of Immunity—The Pathogen's Perspective

- Learning the genome sequences of pathogens can reveal how they infect, which provides clues to developing new treatments.
- Crowd diseases spread rapidly through a population that has had no prior exposure, passed from members of a population that have had time to adapt to the pathogen.
- Throughout history, people have used bacteria and viruses as weapons.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

- Match the cell type to the type of biochemical it produces.

1. mast cell	a. antibodies
2. T cell	b. HLA class II genes
3. B cell	c. interleukin
4. macrophage	d. histamine
5. all cells with nuclei	e. interferon
6. antigen-presenting cell	f. heparin
	g. tumor necrosis factor
	h. HLA class I genes
- What does "nonself" mean? Give an example of a nonself cell in your own body.
- Distinguish between viruses and bacteria.
- What is the physical basis of a blood type?
- Distinguish between using serology or genotyping to type blood.
- Explain why an HLA-disease association is not a diagnosis.
- Explain how mucus, tears, cilia, and ear wax are part of the immune response.
- Distinguish between
 - a T cell and a B cell.
 - innate and adaptive immunity.
 - a primary and secondary immune response.
 - a cellular and humoral immune response.
 - an autoimmune condition and an allergy.
 - an inherited and acquired immune deficiency.
- Which components of the human immune response explain why we experience the same symptoms of an upper respiratory infection (a "cold") when many different types of viruses can cause these conditions?

10. State the function of each of the following immune system biochemicals:
 - a. complement proteins
 - b. collectins
 - c. antibodies
 - d. cytokines
11. What does HIV do to the human immune system?
12. Cite three reasons why developing a vaccine against HIV infection has been challenging.
13. What would be the consequences of lacking
 - a. helper T cells?
 - b. cytotoxic T cells?
 - c. B cells?
 - d. macrophages?
14. Explain how the immune system can respond to millions of different nonself antigens, if there are only a few hundred antibody genes.
15. How are SCID and AIDS similar and different?
16. What part do antibodies play in allergic reactions and in autoimmune disorders?
17. What do a plasma cell and a memory cell descended from the same B cell have in common? How do they differ?
18. Why is a deficiency of T cells more dangerous than a deficiency of B cells?
19. Cite two explanations for why autoimmune disorders are more common in females.
20. How do each of the following illnesses disturb immunity?
 - a. graft-versus-host disease
 - b. SCID
 - c. scleroderma
 - d. AIDS
 - e. hayfever
21. Why is a polyclonal antibody response valuable in the body, but a monoclonal antibody valuable as a diagnostic tool?
22. State how each of the following alters immune system functions:
 - a. a vaccine
 - b. an antibiotic drug
 - c. a cytokine-based drug
 - d. an antihistamine drug
 - e. a transplant

Applied Questions

1. "Winter vomiting disease," a form of gastroenteritis sometimes called "stomach flu," is caused by a virus called norovirus. It makes a person miserable for 1 to 2 days. Why do some people get the illness every year?
2. Rasmussen's encephalitis causes 100 or more seizures a day. Affected children have antibodies that attack brain cell receptors that normally bind neurotransmitters. Is this condition most likely an inherited immune deficiency, an adaptive immune deficiency, an autoimmune disorder, or an allergy? State a reason for your answer.
3. In the TV program *House*, a talented physician and his staff confront difficult-to-diagnose medical cases. They often have to hypothesize whether symptoms are due to an infection, allergy, poison, autoimmunity, or genetic disease. Discuss how these alternatives might be distinguished.
4. In people with a certain HLA genotype, a protein in their joints resembles an antigen on the bacterium that causes Lyme disease. This infection is transmitted in a tick bite and causes flulike symptoms and joint pain (arthritis). When these individuals become infected, their immune systems attack the bacteria and their joints. Explain why antibiotics treat the early phase of the disease, but not the arthritis.
5. A person exposed for the first time to Coxsackie virus develops a painful sore throat. How is the immune system alerted to the exposure to the virus? When the person encounters the virus again, why doesn't she develop symptoms?
6. A young woman who has aplastic anemia will soon die as her lymphocyte levels drop sharply. What type of cytokine might help her?

Web Activities

7. Many websites describe products (food supplements) that supposedly "boost" immune system function. Locate such a website and identify claims that are unclear, deceptive, vague, or incorrect. Alternatively, identify a claim that is consistent with the description of immune system function in this chapter.

Case Studies and Research Results

8. State whether each of the following situations involves an autograft, an isograft, an allograft, or a xenograft.
 - a. A man donates part of his liver to his daughter, who has a liver damaged by cystic fibrosis.
 - b. A woman with infertility receives an ovary transplant from her identical twin.
 - c. A man receives a heart valve from a pig.
 - d. A woman who has had a breast removed has a new breast built using her fatty thigh tissue.
9. Mark and Louise are planning to have their first child, but they are concerned because they think that they have an Rh incompatibility. He is Rh⁻ and she is Rh⁺. Will there be a problem? Why or why not?



Thyroid cancer is often discovered on autopsy. The cancer is slow-growing, and may not produce symptoms. Hence, it is sometimes never noticed. In this person the tumor is obvious as a swelling in the neck.

CHAPTER

18

Genetics of Cancer

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Microarrays Illuminate Thyroid Cancer

I never thought I would care much about the cells of my thyroid gland. That changed on August 4, 1993, when my physician, looking at me from across a room, said, “What’s that lump in your neck?”

Soon after, a specialist stuck six thin needles into the lump to sample thyroid cells for testing, telling me that 99 percent of thyroid “nodules” are not cancerous. However, when he approached with a seventh needle for a sample for a study on something called *p53*, I began to worry, because *p53* is a gene associated with cancer. When the specialist called early on a Monday morning, I knew I was among the unlucky 1 percent. I had papillary thyroid cancer, which accounts for 80 percent of cases and is easily treated with surgery and radioactive iodine. But when I was on the operating table, the surgeon did not think my nodule looked like a papillary tumor. Off it went to the pathology lab, while I waited on the table. The results: I had two tumors, one papillary, one follicular. Treatment was successful.

Had I developed thyroid cancer today, I might not have had to wait on an operating table while a pathologist examined my cells for the telltale distinctions between tumor types. DNA microarrays can now highlight five key genes that are expressed differently in papillary and follicular thyroid cancers. My physicians would have known, before surgery, the genetic nature of my tumors. This approach is very valuable for cancer in which treatment differs depending upon the genetic profiles of the cells, such as breast or prostate cancers.

18.1 Cancer Is Genetic, But Usually Not Inherited

Cancer has been part of human existence for eons. Egyptian mummies from 3000 B.C. show evidence of cancerous tumors, and by 1600 B.C., the Egyptians were attempting to treat cancer. Papyruses illustrate them cutting or burning off growths, and using more inventive treatments for less obvious tumors. A remedy for uterine cancer, for example, introduced fresh ground dates mixed with pig's brain into the vagina!

By 300 B.C., Hippocrates had described several types of tumors, and coined the term “cancer” to describe the crablike shape of a tumor invading normal tissue. He attributed cancer to a buildup of black bile; others blamed it on fermenting lymph, injury, irritation, or simply “melancholia.” Today we know that the collection of diseases called cancer reflects a profound derangement of the cell cycle that can be set into motion by environmental factors. Sequences of mutations in somatic cells and gene expression changes underlie the progression of cancer as it spreads.

Cancer has or will affect one in three of us. Diagnosis and treatment are becoming increasingly individualized, thanks largely to genetic and genomic approaches to describing cancer cells.

Cancer is a complication of being a many-celled organism. Our specialized cells must follow a schedule of mitosis—the cell cycle—so that organs and other body parts either grow appropriately during childhood, stay a particular size and shape in an adult, or repair damage by replacing tissue. If a cell in solid tissue escapes normal controls on its division rate, it forms a growth called a tumor (**figure 18.1**). In the blood, such a cell divides to take over the population of blood cells.

A tumor is benign if it grows in place but does not spread into, or “invade,” surrounding tissue. A tumor is cancerous, or malignant, if it infiltrates nearby tissue. A malignant tumor also sends parts of itself into the bloodstream or lymphatic vessels, either of which transports it to other areas, where the cancer cells “seed” the formation of new tumors. The process of spreading is termed **metastasis**, which means “not standing still.”

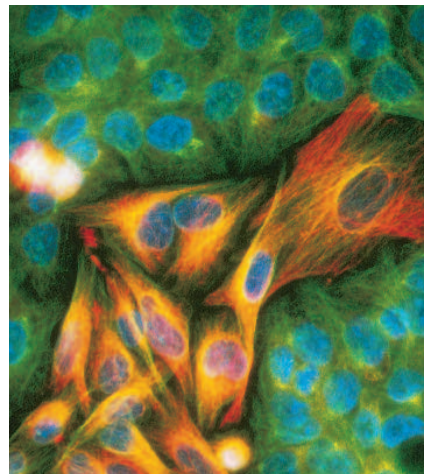
Cancer is a group of disorders that arise from alterations in genes. Only about 10 percent of cases are inherited as single-gene disorders, in which the faulty instructions are in every cell. More often, mutations in cancer-causing genes occur in a few somatic cells over a lifetime. Cancer is usually a genetic disease at the cellular level, but not at the whole-body level.

Combinations of particular gene variants sum to increase the risk of cancer by making cells more sensitive to environmental factors that affect the cell cycle. As a result, cancer may “run in families” yet not follow a single-gene pattern of inheritance. Cancer often takes years to develop, as a sequence of genes mutate in the affected tissue. Then, the cells whose mutations enable them to divide more often than others gradually take over the tissue. Even though a cancer may not spread for years, certain mutations or changes in gene expression can indicate that it will.

Cancer wasn't always considered a genetic phenomenon. When President Richard Nixon declared a “war on cancer” in 1971, the targets were radiation, viruses, and chemicals. These



a.



b.

Figure 18.1 Cancer cells stand out. (a) A melanoma is a cancer of the pigment-producing cells (melanocytes) in the skin. It may have any or all of four characteristics, abbreviated ABCD: it is **a**symmetric, has **b**orders that are irregular, **c**olor variations, and a **d**iameter of more than 5 millimeters. These tumors were on a woman's back. **(b)** Stains and dyes reveal cancer at the cellular level. These melanoma cells stain orange. The different staining characteristics of cancer cells reflect differences in gene expression patterns between the normal and cancerous states.

agents actually cause cancer by interfering with the precise genetic controls of cell division.

An early hint at the genetic nature of cancer was the observation that most substances known to be carcinogens (causing cancer) are also mutagens (damaging DNA). Researchers first discovered genes that could cause cancer in 1976. These **oncogenes** were originally identified in viruses that cause cancer in rodents. Then the genes' normal equivalents, which control the cell cycle, were found, in rodents and humans.

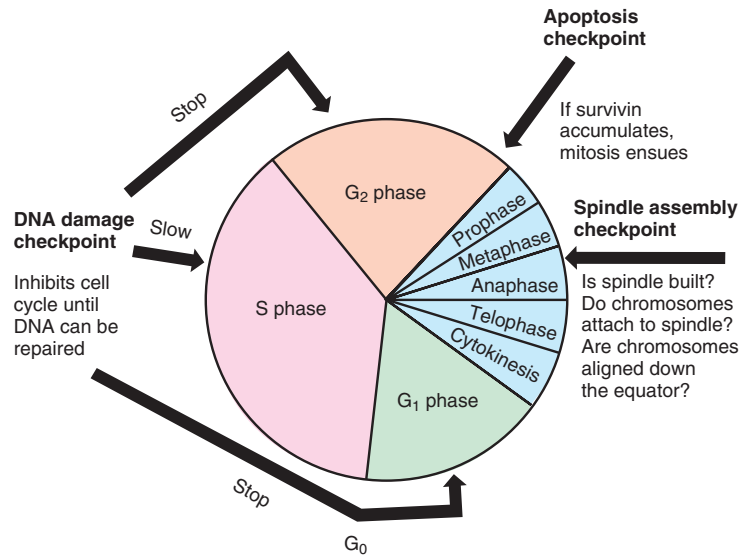


Figure 18.2 Cell cycle checkpoints. Checkpoints ensure that mitotic events occur in the correct sequence. Many types of cancer result from faulty checkpoints.

In the 1980s and 1990s, searches for cancer-causing genes began with rare families that had many young members who had the same type of cancer and specific unusual chromosomes. Then the search focused on genes in the identified chromosome region whose protein products could affect cell cycle control, leading to the discovery of more than 100 oncogenes. An oncogene is a gene that causes cancer when inappropriately activated. Family studies also identified more than 30 **tumor suppressor** genes, which cause cancer when they are deleted or inactivated. The normal function of a tumor suppressor gene is to keep the cell cycle running at the appropriate rate for a particular cell type under particular conditions.

In addition to activated oncogenes and inactivated tumor suppressor genes, changes in gene expression accompany cancer. DNA microarrays highlight mutations and patterns of gene expression that paint “molecular portraits” of the disease. These views are making it possible to recognize subtypes of cancers affecting the same cell types, as the opener to chapter 11 discusses for leukemia. The subtypes explained why the prevailing treatment did not work for some young patients.

Loss of Cell Cycle Control

Cancer is a consequence of cell cycle disruption. **Figure 18.2** repeats the cell cycle diagram from chapter 2. Cancer begins when a cell divides more frequently, or more times, than the noncancerous cell it descended from (**figure 18.3**).

The timing, rate, and number of mitoses a cell undergoes depend on protein growth factors and signaling molecules from outside the cell, and on transcription factors from within. Because these biochemicals are under genetic control, so is the cell cycle. Cancer cells probably arise often, because mitoses are so frequent that an occasional cell escapes control. However, the immune system destroys most cancer cells after recognizing tumor-specific antigens on their surfaces.

The discovery of the checkpoints that control the cell cycle revealed how cancer can begin. A mutation in a gene that normally halts or slows the cell cycle can lift the constraint, leading to inappropriate mitosis. Failure to pause long enough to repair DNA can allow a mutation in an oncogene or tumor suppressor gene to persist.

Loss of control over telomere length may also contribute to cancer by affecting the cell cycle. Recall that telomeres, or chromosome tips, protect chromosomes from breaking. Human telomeres consist of the DNA sequence TTAGGG repeated thousands of times. The repeats are normally lost from the telomere ends as a cell matures, from 15 to 40 nucleotides per cell division. The more specialized a cell, the shorter its telomeres. The chromosomes in skin, nerve, and muscle cells, for example, have short telomeres. Chromosomes in a sperm cell or oocyte, however, have long telomeres. This makes sense—as the precursors of a new organism, gametes must retain the capacity to divide many times.

Gametes keep their telomeres long thanks to an enzyme, telomerase, that is a complex of RNA and protein. Part of the RNA—the sequence AAUCCC—serves as a template for the 6-DNA-base repeat TTAGGG that builds telomeres. Telomerase moves down the DNA like a zipper, adding six “teeth” (bases) at a time.

In normal, specialized cells, telomerase is turned off, and telomeres shrink, signaling a halt to cell division when they reach a certain size. In cancer cells, telomerase is turned back on. Telomeres extend, and this releases the normal brake on rapid cell division. As daughter cells of the original abnormal cell continue to divide uncontrollably, a tumor forms, grows, and may spread. Usually the longer the telomeres in cancer cells, the more advanced the disease. However, turning on telomerase production in a cell is not sufficient in itself to cause cancer. Many other things must go wrong for cancer to begin.

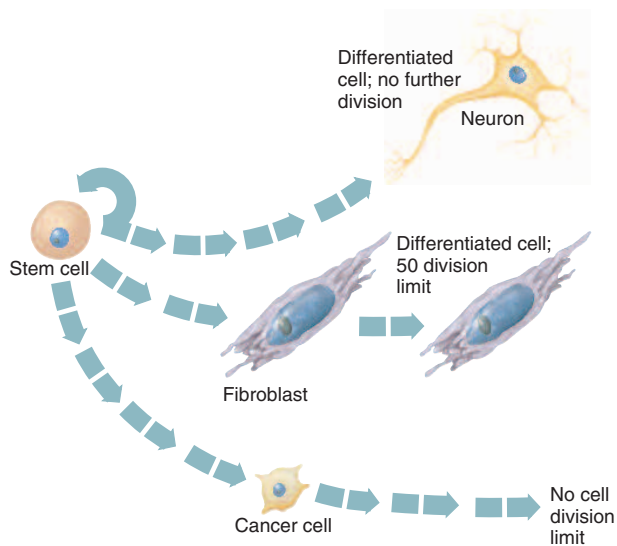


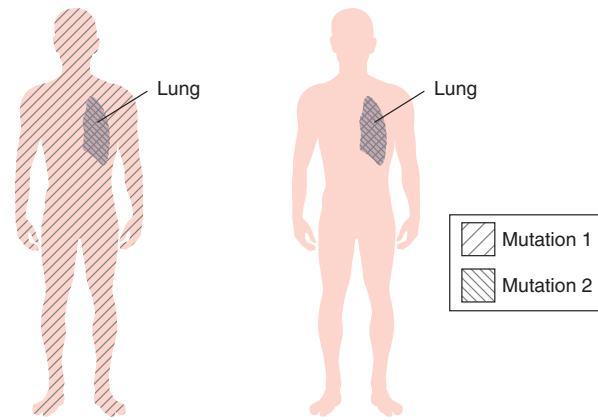
Figure 18.3 Cancer sends a cell down a pathway of unrestricted cell division. Cells may be terminally differentiated and no longer divide, such as a neuron, or differentiated yet still capable of limited cell division, such as a fibroblast (connective tissue cell). Cancer cells either lose specializations or never specialize; they divide unceasingly. (Arrows represent some cell divisions; not all daughter cells are shown.)

Inherited Versus Sporadic Cancer

Although cancer genes were discovered in families with inherited cancers, most cancers are sporadic, which means that the causative mutations occur only in cells of the affected tissue. These are **somatic mutations** because they occur in nonsex cells. A sporadic cancer may result from a single dominant mutation or from two recessive mutations in the same gene. The cell harboring the mutation loses control of its cell cycle, divides continuously, and a tumor forms and grows.

Susceptibility to developing a sporadic cancer is *not* directly passed on to future generations because the gametes do not carry the mutant allele or alleles. In contrast are **germline mutations**, in which cancer susceptibility *is* directly passed to future generations because the mutations are in every cell, including gametes. Cancer develops when a second mutation occurs in the other allele in a somatic cell in the affected body part (**figure 18.4**).

Germline mutations may explain why some heavy smokers develop lung cancer, but many do not; the unlucky ones may have inherited a susceptibility allele in every cell. Years of exposing lung tissue to the carcinogens in smoke eventually cause a mutation in a tumor suppressor gene or oncogene of a lung cell, giving it a proliferative advantage. Without the susceptibility gene, two such somatic mutations are necessary to trigger the cancer. This, too, can be the result of an environmental insult, but it takes longer for two events to occur than one. Germline cancers are rare, but they have high penetrance and tend to strike earlier in life than sporadic cancers.



a. Germline (inherited) cancer b. Sporadic cancer

Figure 18.4 Germline versus sporadic cancer. (a) In germline cancer, every cell has one gene variant that increases cancer susceptibility, and a second mutation in a cell of the affected tissue. This type of predisposition to cancer is inherited as a single-gene trait. (b) A sporadic cancer forms when a dominant mutation occurs in a somatic cell or two recessive mutations occur in the same gene in the same somatic cell. An environmental factor can cause the somatic mutations of cancer. Note that each lung has undergone both mutations 1 and 2.

Key Concepts

1. Cancer is genetic, but not usually inherited.
2. Single genes (oncogenes and tumor suppressors), when mutant, can cause cancer. Cancer cells have different gene expression profiles compared to the cells from which they descend.
3. Cancer is caused by a loss of cell division control. Implicated genes encode growth factors, transcription factors, or telomerase.
4. Most cancer mutations occur in somatic cells.
5. Cancer may develop when an environmental trigger mutates a somatic cell (sporadic) or when a somatic mutation compounds an inherited susceptibility (germline).

18.2 Characteristics of Cancer Cells

Cancer begins at the genetic and cellular levels. If not halted, it spreads through tissues to take over organs and organ systems. **Figure 18.5** summarizes the steps in the origin and spread of a cancer.

Cell division is rigorously controlled. Whether a cell divides or stops dividing and whether it differentiates depends upon signals from surrounding cells. A cancer cell simply stops “listening” to those signals.

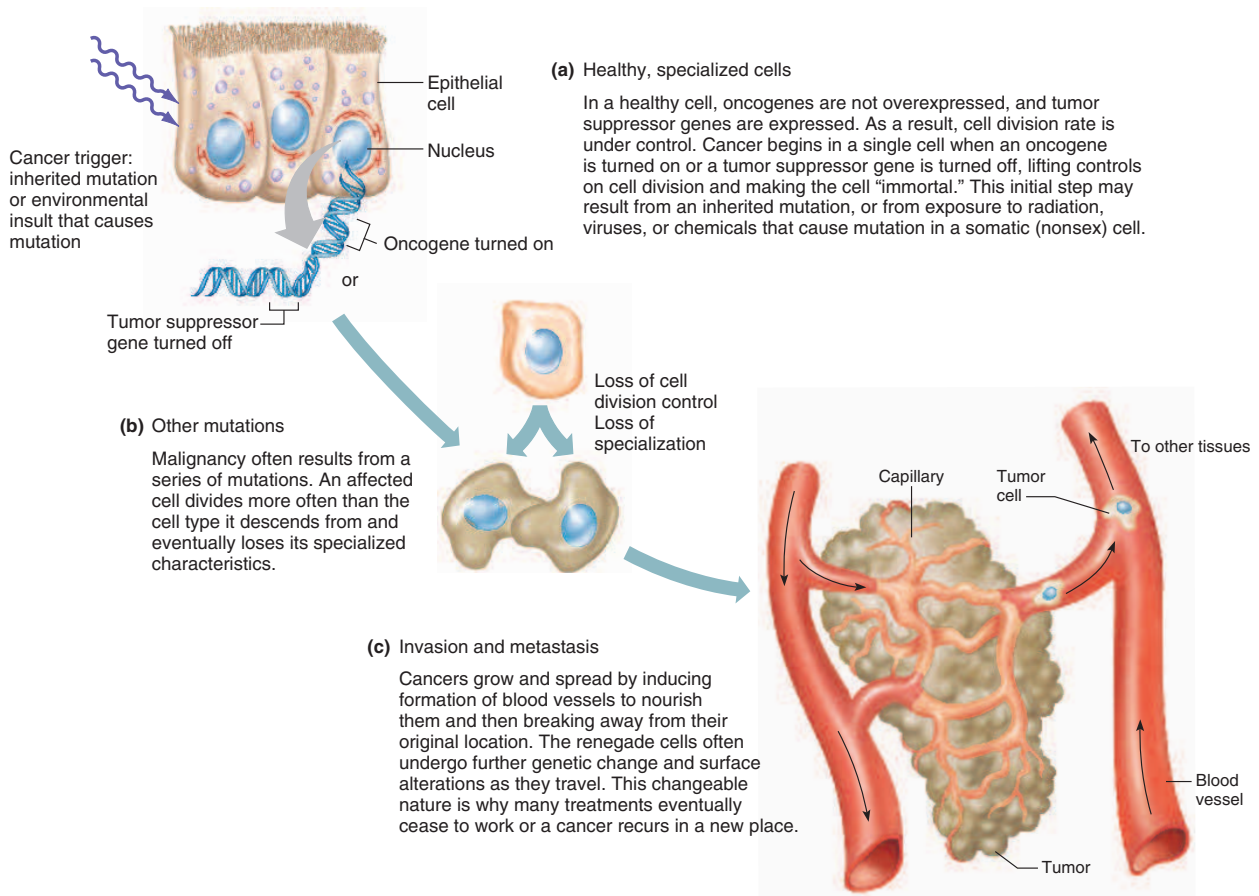


Figure 18.5 Steps in the development of cancer.

Cancer cells can divide continuously if given sufficient nutrients and space. Cervical cancer cells of a woman named Henrietta Lacks, who died in 1951, vividly illustrate the hardiness of these cells. Her cells persist today as standard cultures in many research laboratories. These “HeLa” cells divide so vigorously that when they contaminate cultures of other cells they soon take over.

Cells vary greatly in their capacity to divide. Cancer cells divide more frequently or more times than the cells from which they arise. Yet even the fastest-dividing cancer cells, which complete mitosis every 18 to 24 hours, do not divide as often as some cells in a normal human embryo do. Still, some cancers grow alarmingly fast. The smallest detectable fast-growing tumor is half a centimeter in diameter and can contain a billion cells. These cells divide, producing a million or so new cells in an hour. If 99 percent of the tumor’s cells are destroyed, 10 million are left to proliferate. Other cancers develop over years. A tumor grows more slowly at first because fewer cells divide. By the time the tumor is the size of a pea—when it is usually detectable—billions of cells are actively dividing. A cancerous tumor eventually grows faster than surrounding tissue because a greater proportion of its cells is dividing.

A cancer cell looks different from a normal cell. It is rounder because it does not adhere to surrounding normal cells

as strongly as other cells do. Because the plasma membrane is more fluid, different substances cross it. A cancer cell’s surface may sport different antigens than are on other cells or different numbers of antigens that are also on normal cells. The “prostate specific antigen” (PSA) blood test that indicates increased risk of prostate cancer, for example, detects elevated levels of this protein that may come from cancer cells.

When a cancer cell divides, both daughter cells are cancerous, since they inherit the altered cell cycle control. Therefore, cancer is said to be heritable because it is passed from parent cell to daughter cell. A cancer is also transplantable. If a cancer cell is injected into a healthy animal of the same species, it will proliferate there.

A cancer cell is **dedifferentiated**, which means that it is less specialized than the normal cell types near it that it might have descended from. A skin cancer cell, for example, is rounder and softer than the flattened, scaly, healthy skin cells above it in the epidermis, and is more like a stem cell in both appearance and division rate. Cancer cell growth is unusual. Normal cells in a container divide to form a single layer; cancer cells pile up on one another. In an organism, this pileup would produce a tumor. Cancer cells that grow all over one another are said to lack contact inhibition—they do not stop dividing when they crowd other cells.

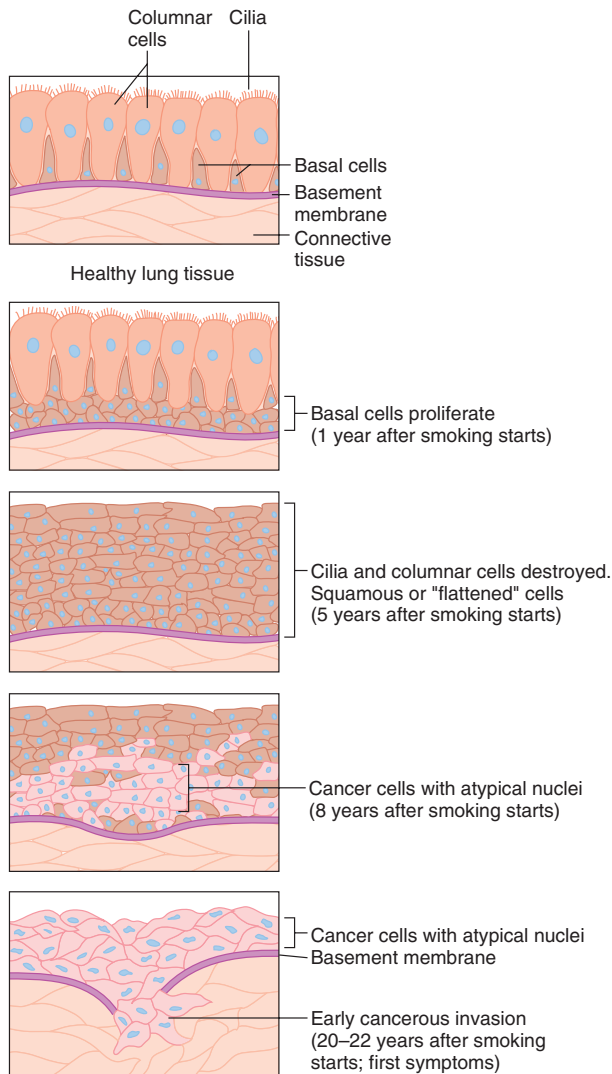


Figure 18.6 Some cancers take years to spread. Lung cancer due to smoking begins with irritation of the lining tissue in respiratory tubes. Ciliated cells die and basal cells divide. If the irritation continues, cancerous changes occur.

Cancer cells have surface structures that enable them to squeeze into any space, a property called **invasiveness** (figure 18.6). They anchor themselves to tissue boundaries, called basement membranes, where they secrete enzymes that cut paths through healthy tissue. Unlike a benign tumor, an invasive malignant tumor grows irregularly, sending tentacles in all directions.

Cancer cells eventually reach the bloodstream or lymphatic vessels, which take them to other parts of the body—unless treatment stops the disease process. The traveling cancer cells settle into new sites—this is metastasis.

Once a tumor has grown to the size of a pinhead, interior cancer cells respond to the oxygen-poor environment by

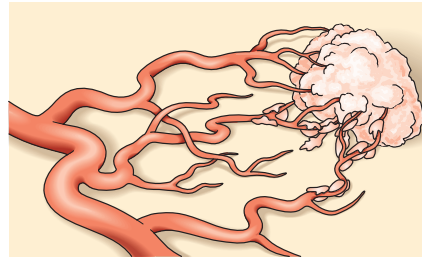


Figure 18.7 Angiogenesis nurtures a tumor. Cells starved for oxygen deep within a tumor secrete vascular endothelial growth factor (VEGF), which stimulates nearby capillaries to extend branches toward a tumor. A class of drugs treats cancer by blocking VEGF.

secreting a protein, called vascular endothelial growth factor (VEGF). It stimulates nearby capillaries (the tiniest blood vessels) to sprout new branches that extend toward the tumor, bringing in oxygen and nutrients and removing wastes. This growth of new capillary extensions is called **angiogenesis**, and it is critical to a cancer's growth and spread. Capillaries may snake into and out of the tumor (figure 18.7). Cancer cells wrap around the blood vessels and creep out upon this scaffolding, invading nearby tissue. In addition to attracting their own blood supply, cancer cells may also secrete hormones that encourage their own growth. This is a new ability because the cells they descend from do not produce these hormones.

Once cancer cells move to a new body part, the disease has metastasized. The DNA of secondary tumor cells often mutates, and chromosomes may break or rearrange, as DNA in primary tumors does too. Many cancer cells are aneuploid. The metastasized cancer thus becomes a new genetic entity that may resist treatments that were effective against most cells of the original tumor. Because gene expression patterns associated with metastasis are detectable early, new cancer treatments may be designed to prevent metastasis.

Table 18.1 summarizes the characteristics of cancer cells. **Table 18.2** lists the processes and pathways that can be abnormal in cancer. The many mutations that cause or contribute to cancer converge in ways that affect these few pathways.

Key Concepts

1. Cancer occurs when cells divide faster or more times than normal.
2. Cancer cells are heritable, transplantable, and dedifferentiated. They lack contact inhibition, cutting through basement membranes.
3. A cancerous growth is invasive and can metastasize and stimulate angiogenesis, spreading farther.

Table 18.1 Characteristics of Cancer Cells

Oilier, less adherent
Loss of cell cycle control
Heritable
Transplantable
Dedifferentiated
Lack contact inhibition
Induce local blood vessel formation (angiogenesis)
Invasive
Increased mutation rate
Can spread (metastasize)

Table 18.2 Processes and Pathways Affected in Cancer

	Discussed in Chapter
Angiogenesis	18
Apoptosis	7
Cell adhesion	2
Cell cycle control	2
DNA repair	12
Metabolism of carcinogens	18
Signal transduction	2

18.3 Origins of Cancer Cells

Mutations that turn a cell cancerous are only a first step in the disease process. Factors that influence whether or not cancer develops include how specialized the initial cell is and the location of that cell in the tissue.

Cancer can begin at a cellular level in at least four ways:

- activation of stem cells that produce cancer cells
- dedifferentiation
- increase in the proportion of a tissue that consists of stem or progenitor cells
- faulty tissue repair.

Most cancer cells are more specialized than stem cells, but considerably less specialized than the differentiated cells near them in a tissue (see figures 2.23 and 2.25). From which does the cancer cell arise? A cancer cell may descend from a stem cell that yields slightly differentiated daughter cells that

retain the capacity to self-renew, or a cancer cell may arise from a specialized cell that loses some of its features and can divide. Certain stem cells, called **cancer stem cells**, veer from normal development and produce both cancer cells and abnormal specialized cells. Cancer stem cells are found in cancers of the brain, blood, and epithelium (particularly in the breast, colon, and prostate).

Figure 18.8 illustrates how cancer stem cells may cause brain tumors. In (a), as cancer stem cells give rise to progenitors and then differentiated cells (neurons, astrocytes, and oligodendrocytes), a cell surface molecule called CD133 is normally lost (designated CD133⁻) at the late progenitor stage. In contrast, in (b), cancer cells retain the molecule (designated CD133⁺). Some progenitor cells that descend from a cancer stem cell can relentlessly divide, and they ultimately accumulate, forming a brain tumor.

Another route to cancer may be cells that lose some of their distinguishing characteristics as mutations occur when they divide. Or, cells on the road to cancer may begin to express “stemness” genes that override signals to remain specialized (**figure 18.9**). Whatever the mechanism, the result is dedifferentiation. So far experiments have not captured the exact moment when a cell both loses specializations and becomes able to continually divide. However, researchers have identified a biochemical, named “reversine,” that can stimulate differentiated cells to divide and give rise to progenitor cells in mice. Reversine may play a role in the dedifferentiation of cancer cells.

Another possible origin of cancer may be a loss of balance at the tissue level in favor of cells that can divide continually or frequently—like a population growing faster if more of its members are of reproductive age. Consider a tissue that is 5 percent stem cells, 10 percent progenitors, and 85 percent differentiated cells. If a mutation, over time, shifts the balance in a way that creates more stem and progenitor cells, the extra cells pile up, and a tumor forms (**figure 18.10**).

Uncontrolled tissue repair may cause cancer (**figure 18.11**). If too many cells divide to fill in the space left by injured tissue, and those cells keep dividing, an abnormal growth may result.

With so many millions of cells undergoing so many error-prone DNA replications, and so many ways that cancer can arise, it perhaps isn’t surprising that cancer is so common. Yet most of the time, the immune system vanquishes a cancer before it progresses very far.

Key Concepts

1. Cancer stem cells produce cancer cells and abnormal specialized cells.
2. Dedifferentiation might occur through mutation or overexpression of “stemness” genes.
3. Upsetting the balance of stem and progenitor to differentiated cells can cause cancer as excess, fast-dividing cells accumulate.

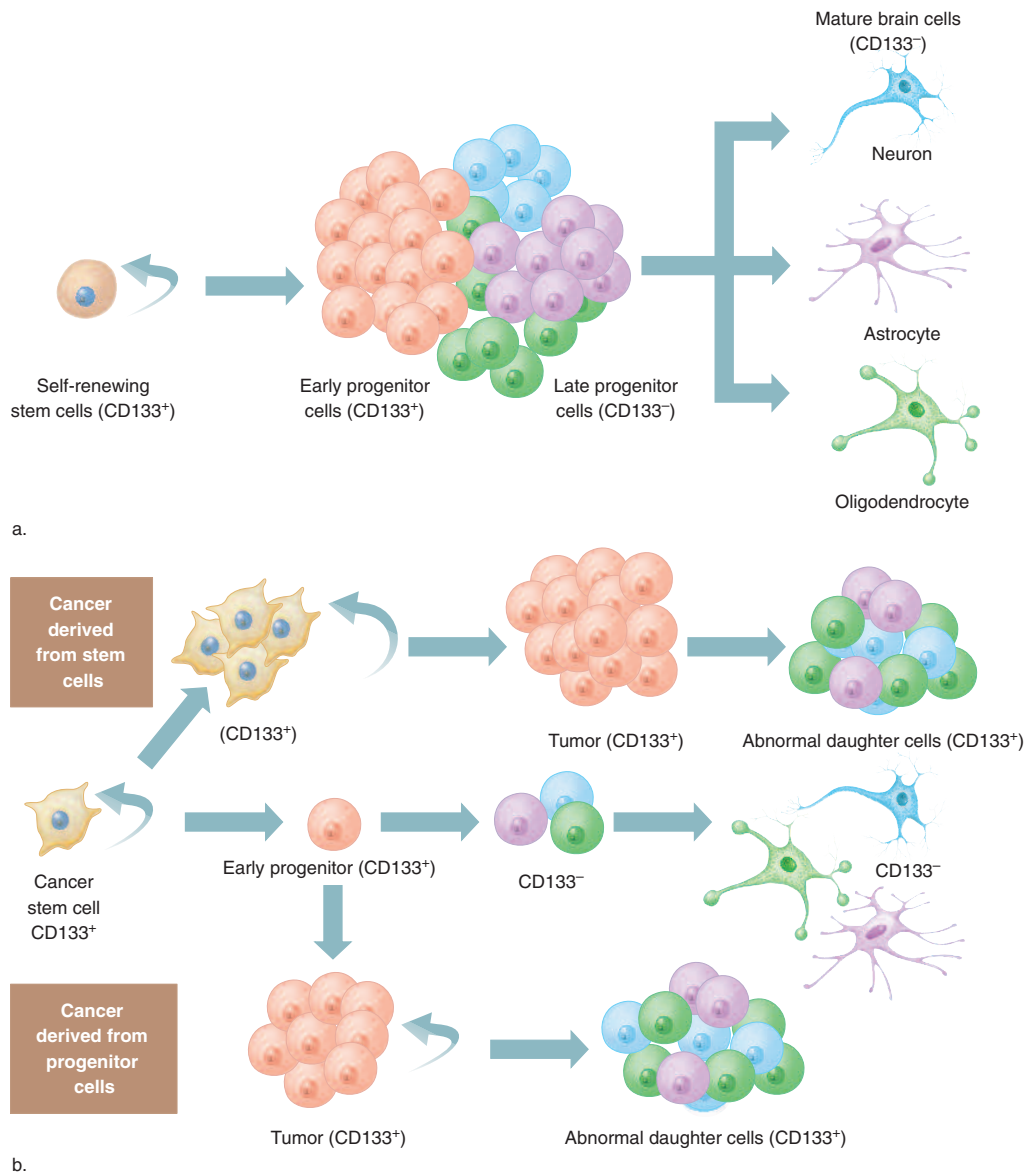


Figure 18.8 Cancer stem cells. (a) In the developing brain, stem cells divide to self-renew and give rise to early progenitor cells, which in turn divide to yield late progenitor cells. These late progenitor cells lose the CD133 cell surface marker, and they divide to give rise to daughter cells that specialize as neurons or two types of supportive cells, astrocytes or oligodendrocytes. (b) A cancer stem cell can divide to self-renew and give rise to a cancer cell, which in turn can also spawn abnormal daughter cells. Some early progenitors give rise to normal differentiated cells. Sometimes the cancer-causing mutations occur in the cancer stem cell-derived early progenitor cell. In this case, the early progenitors form the tumor, which may spawn some abnormal daughter cells. Note that stem cells, cancer stem cells, early progenitor cells, and abnormal daughter cells all have the CD133⁺ marker, but the differentiated cells do not.

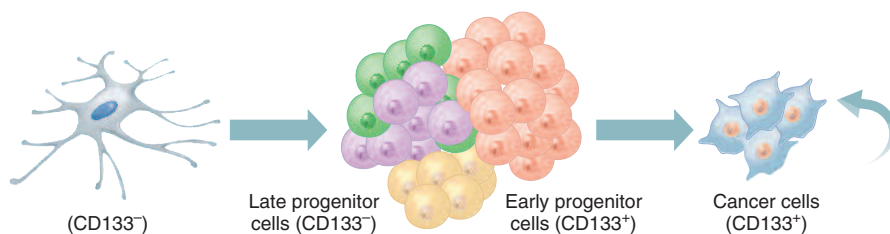


Figure 18.9 Dedifferentiation reverses specialization. Mutations in a differentiated cell could reactivate latent “stemness” genes, giving the cell greater capacity to divide while causing it to lose some of its specializations. These are two of the defining characteristics of cancer cells.

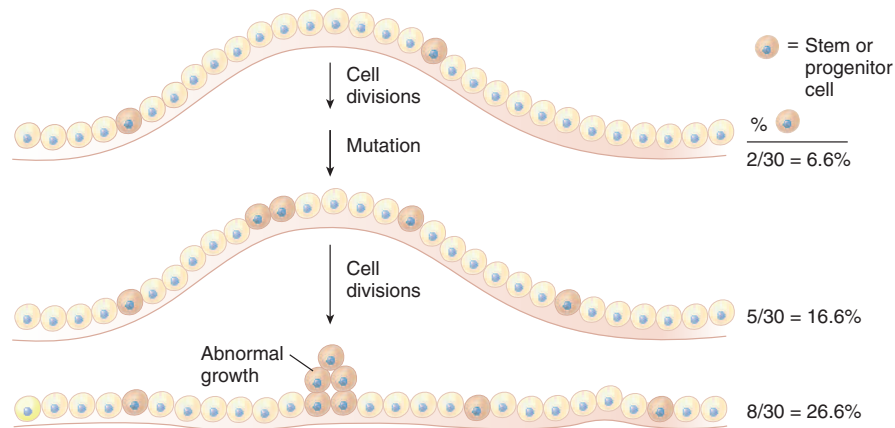


Figure 18.10 Shifting the balance in a tissue toward cells that divide. If a mutation renders a differentiated cell able to divide to yield other cells that frequently divide, then over time these cells may take over, forming an abnormal growth.

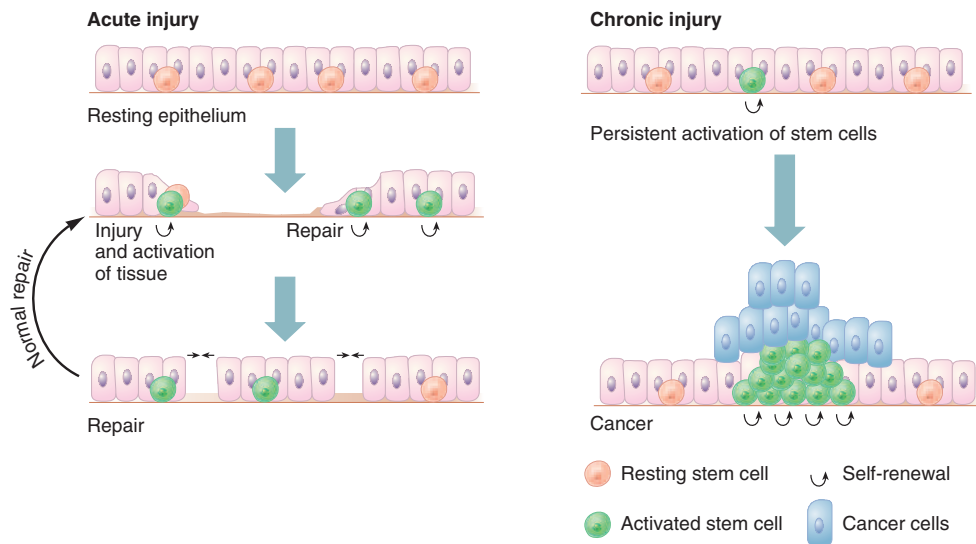


Figure 18.11 Too much repair may trigger tumor formation. If epithelium is occasionally damaged, resting stem cells can become activated and divide to fill in the tissue. If injury is chronic, the persistent activation of stem cells to renew the tissue can veer out of control, fueling an abnormal growth.

18.4 Cancer Genes and MicroRNAs

Most mutations that cause cancer are in oncogenes or tumor suppressor genes. A third category includes mismatch mutations in DNA repair genes (see section 12.6) that allow other mutations to persist. When such mutations activate oncogenes or inactivate tumor suppressor genes, cancer results. DNA repair disorders are often inherited in a single-gene fashion, and are quite rare. They tend to cause diverse and widespread tumors.

Oncogenes

Genes that normally trigger cell division when it is appropriate are called **proto-oncogenes**. They are active where and when high rates of cell division are necessary, such as in a

wound or in an embryo. When proto-oncogenes are turned on at the wrong time or place, they function as oncogenes (“onco” means cancer). Usually oncogene activation is associated with a point mutation, chromosomal translocation, or inversion, and a gain of function. In contrast, a tumor suppressor gene mutation that causes cancer is usually a deletion that removes a function.

Abnormal activation of a proto-oncogene into an oncogene may be the result of either a mutation or a change in expression of the wild type gene. A single base change in a proto-oncogene causes bladder cancer, for example. Alternatively, a proto-oncogene may be moved near a gene that is highly expressed; then it, too, is rapidly or frequently transcribed. A human proto-oncogene is normally activated in cells at the site of a wound, where it stimulates production

of growth factors that cause mitosis to fill in the damaged area with new cells. When that proto-oncogene is activated at a site other than a wound—as an oncogene—it still hikes growth factor production and stimulates mitosis. However, because the site of the action is not damaged tissue, the new cells form a tumor.

Some proto-oncogenes encode transcription factors that, as oncogenes, are too highly expressed. (Recall from chapter 10 that transcription factors bind to specific genes and activate transcription.) The products of these activated genes then contribute to the cancer cell's characteristics. Oncogenes may also block apoptosis. As a result, damaged cells do not die, but divide.

Increased Expression in a New Location

A proto-oncogene can become an oncogene when it is placed next to a gene that boosts its expression. A virus infecting a cell, for example, may insert DNA next to a proto-oncogene. When the viral DNA is rapidly transcribed, the adjacent proto-oncogene (now an oncogene) is also rapidly transcribed. Increased production of the oncogene's encoded protein then switches on genes that promote mitosis, triggering the cascade of changes that leads to cancer. Viruses cause cervical cancer, Kaposi sarcoma, and acute T cell leukemia.

A proto-oncogene can also be activated when it is moved next to a gene that is normally very actively transcribed. This can happen when a chromosome is inverted or translocated, placing a gene in a new chromosomal environment. For example, a cancer of the parathyroid glands in the neck is associated with an inversion on chromosome 11, which places a proto-oncogene next to a DNA sequence that controls transcription of the parathyroid hormone gene. When the gland synthesizes the hormone, the oncogene is expressed, too. Cells in the gland divide, forming a tumor.

Ironically, the immune system contributes to cancer when a translocation or inversion places a proto-oncogene next to an

antibody gene. Recall from chapter 17 that antibody genes normally move into novel combinations when a B cell is stimulated and they are very actively transcribed. Cancers associated with viral infections, such as cervical cancer following HPV infection, may be caused when proto-oncogenes are mistakenly activated with antibody genes. Similarly, in Burkitt lymphoma, a cancer common in Africa, a large tumor develops from lymph glands near the jaw. People with Burkitt lymphoma are infected with the Epstein-Barr virus, which stimulates specific chromosome movements in maturing B cells to assemble antibodies against the virus. A translocation places a proto-oncogene on chromosome 8 next to an antibody gene on chromosome 14. The oncogene is overexpressed, and the cell division rate increases. Tumor cells of Burkitt lymphoma patients have the translocation (**figure 18.12**).

We can use the information of changes in gene expression that promote cancer to diagnose, treat, or track response to treatment, even without knowing what the expression patterns mean. For example, ocular melanoma affects pigment cells in the eye—it is much more deadly than common skin melanoma. Many cases of ocular melanoma spread, 95 percent to the liver. Researchers extracted mRNAs from affected eyes and measured the expression of ten genes. They derived two “molecular signatures”—patterns of mRNAs that are more or less abundant than normal—one predicting a low risk of spread to the liver, the other a high risk. This information is used to guide treatment choices.

Fusion Proteins with New Functions

An oncogene is also activated when a proto-oncogene moves next to another gene, and the gene pair is transcribed and translated together, as if they are one gene. The double gene product, called a **fusion protein**, activates or lifts control of cell division.

A fusion oncoprotein causes acute promyelocytic leukemia. (Leukemias differ by the type of white blood cell affected.) A translocation between chromosomes 15 and 17

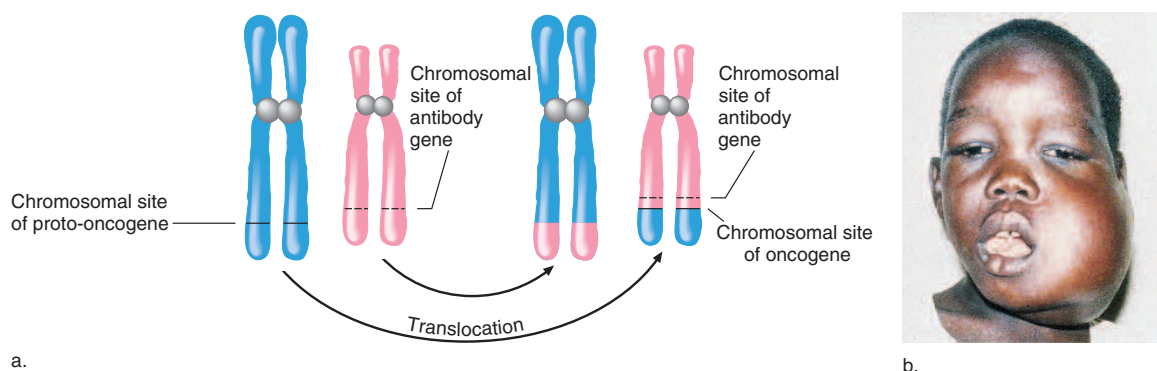


Figure 18.12 A translocation that causes cancer. **(a)** The cause of Burkitt lymphoma is translocation of a proto-oncogene on chromosome 8 to chromosome 14, next to a highly expressed antibody gene. Overexpression of the translocated proto-oncogene, now an oncogene, triggers the molecular and cellular changes of cancer. **(b)** Burkitt lymphoma often affects the jaw.

brings together a gene coding for the retinoic acid cell surface receptor and an oncogene called *myl*. The fusion protein functions as a transcription factor, which, when overexpressed, causes cancer. The nature of this fusion protein explains why some patients who receive retinoid (vitamin A-based) drugs recover. Their immature, dedifferentiated cancer cells, apparently stuck in an early stage of development where they divide frequently, suddenly differentiate, mature, and then die. Perhaps the cancer-causing fusion protein prevents affected white blood cells from getting enough retinoids to specialize, locking them in an embryoniclike, rapidly dividing state. Supplying extra retinoids allows the cells to resume their normal developmental pathway. **Reading 18.1** tells the story of a young magazine editor who recovered from a different type of leukemia thanks to a drug developed from understanding how a fusion protein causes cancer.

Receiving a Too-Strong Division Signal

In about 25 percent of women with breast cancer, affected cells have 1 to 2 million copies of a cell surface protein called *Her-2/neu* that is the product of an oncogene. The normal number of these proteins is only 20,000 to 100,000.

The *Her-2/neu* proteins are receptors for epidermal growth factor. The receptors traverse the plasma membrane, extending outside the cell into the extracellular matrix and also dipping into the cytoplasm. They function as a tyrosine kinase, as is the case for the leukemia described in Reading 18.1. When the growth factor binds to the tyrosine of the receptor, the tyrosine picks up a phosphate group, which signals the cell to activate transcription of genes that stimulate cell division. In *Her-2/neu* breast cancer, too many tyrosine kinase receptors send too many signals to divide.

Her-2/neu breast cancer usually strikes early in adulthood and spreads quickly. However, a monoclonal antibody-based drug called Herceptin binds to the receptors, blocking the signal to divide (see figure 17.16). Interestingly, Herceptin works when the extra receptors arise from multiple copies of the gene, rather than from extra transcription of a single *Her-2/neu* gene.

Tumor Suppressors

Some cancers result from loss or silencing of a gene that normally suppresses tumor formation by blocking the activities of other genes. Such a tumor suppressor gene normally inhibits expression of genes involved in all of the activities that turn a cell cancerous, listed in table 18.2. Cancer can result when a tumor suppressor's control is lifted. This can happen if the gene has a deletion, or if the promoter region binds too many methyl (CH_3) groups, which blocks transcription. Binding of CH_3 groups to "CpG islands"—regions in the starts of genes where the sequence "CG" repeats

many times—turns off transcription. Such hypermethylation is an epigenetic change, because the mRNA sequence is unaffected.

Wilms' tumor is an example of a cancer that develops from loss of tumor suppression. A gene that normally halts mitosis in the rapidly developing kidney tubules in the fetus is absent. As a result, an affected child's kidney retains pockets of cells dividing as frequently as if they were still in the fetus, forming a tumor. Following are descriptions of specific tumor suppressor genes.

Retinoblastoma (RB)

RB (MIM 180200) is a rare childhood eye tumor. In 1597, a Dutch anatomist described the eye cancer as a growth "the size of two fists." In 1886, researchers identified inherited cases. At that time, the only treatment was removal of the affected eye. Today, children with an affected parent or sibling, who have a 50 percent chance of having inherited the mutant *RB* gene, can be monitored from birth so that noninvasive treatment can begin early. Full recovery is common. Often the first abnormal sign is an unusual gray area that appears in an eye in a photograph—the tumor reflects light differently than unaffected parts of the eye.

About half of the 1 in 20,000 infants who develop RB inherit susceptibility to the disorder: They harbor one germline mutant allele for the *RB* gene in each of their cells. Cancer develops in any somatic cell where the second copy of the *RB* gene mutates. Therefore, inherited retinoblastoma requires two point mutations or deletions, one germline and one somatic. In sporadic (noninherited) cases, two somatic mutations occur in the *RB* gene. Either way, RB usually starts in a cone cell of the retina, which provides color vision. Study of RB was the origin of the "two-hit" hypothesis of cancer causation—that two mutations (germline and somatic or two somatic) are required to cause a cancer related to tumor suppressor deletion or malfunction.

Many children with RB have deletions in the same region of the long arm of chromosome 13, which led researchers to the cancer-causing gene. In 1987, they found the *RB* gene and identified its protein product, which linked the cancer to control of the cell cycle. The RB protein normally binds transcription factors so that they cannot activate genes that carry out mitosis. It normally halts the cell cycle at G_1 . When the *RB* gene is mutant or missing, the hold on the transcription factor is released, and cell division ensues.

Mutations in the *RB* gene cause other cancers. Children successfully treated for retinoblastoma often develop bone cancer as teens or bladder cancer as adults. Mutant *RB* genes have been found in the cells of patients with breast, lung, or prostate cancers, or acute myeloid leukemia, who never had the eye tumors. These other cancers may be caused by expression of the same genetic defect in different tissues.

Reading 18.1

Erin's Story: How Gleevec Treats Leukemia

When 23-year-old *Glamour* magazine editor Erin Zammett Ruddy went for a routine physical in November 2001, she expected reassurance that her healthy lifestyle had indeed been keeping her healthy (**figure 1**). After all, she felt great. What she got, a few days later, was a shock. Instead of having 4,000 to 10,000 white blood cells per milliliter of blood, she had more than 10 times that number—and many of the cells were cancerous.

"I had just returned from a nice, long lunch to find a message from my doctor. Could I call back? Something had come up in my blood work," recalled Erin. "I was diagnosed with chronic myelogenous leukemia. CML is cancer, and until very recently, it proved fatal in the vast majority of cases."

Although there is hardly a "good" time to find out that you have cancer, Erin's diagnosis came just a few months after a landmark report of a new drug—and, ironically, an article in *Glamour* about three CML survivors. A successful cancer drug typically helps about 20 percent of the patients who take it, often just extending life a few months. But cancer in the blood had vanished in 53 of 54 initial patients, usually quickly. So Erin contacted the lead researcher, Brian Druker, and joined the group. Her cancer was reversed—with just a pill a day, and no side effects.

The drug, Gleevec, is now the standard treatment for CML and a few other cancers. The story of its development illustrates how understanding the genetic events that start and propel a cancer can guide development of an effective weapon.

The tale of Gleevec began on August 13, 1958, when two men entered hospitals in Philadelphia and reported weeks of fatigue. Each had very high white blood cell counts and were diagnosed with CML. Too many immature white blood cells were crowding the healthy cells. The men's blood samples eventually fell into the hands of pathologist Peter Nowell and cytogeneticist David Hungerford. They had developed ways to stimulate white blood cells to divide in culture, and they probed the chromosomes of both leukemic and normal-appearing white blood cells in the two tired men and five others with CML.



Figure 1 "My third bone marrow biopsy—you never get used to the pain," said Erin. Gleevec has treated her leukemia.

Nowell and Hungerford discovered a small, unusual chromosome that was only in the leukemic cells. This was the first chromosome abnormality to be linked to cancer. Later, it would be dubbed "the Philadelphia chromosome" (Ph¹). The link between the cancer and the chromosome anomaly held up in other patients.

With refinements in chromosome banding, important details emerged. In 1972, Janet Rowley at the University of Chicago used new stains that distinguished AT-rich from GC-rich chromosome regions to tell that Ph¹ is the result of a translocation (see figure 13.19). By 1984, researchers had homed in on the two genes juxtaposed in the translocation between chromosomes 9 and 22. Therein lay the clues that would lead to Erin's treatment.

One gene from chromosome 9 is called the Abelson oncogene (*abl*), and the other gene, from chromosome 22, is called the breakpoint cluster region (*bcr*). Two different fusion genes form. The *bcr-abl* fusion gene is part of the Philadelphia chromosome, and it causes CML. The encoded fusion protein, called the BCR-ABL oncoprotein, is a form of the enzyme tyrosine kinase, which is the normal product of the *abl* gene. The cancer-causing form of tyrosine kinase is active for too long, which sends signals into the cell, stimulating it to divide too many times. (The other fusion gene does not affect health.)

The discovery that a fusion oncoprotein started the cellular changes that cause CML gave drug researchers a target. Through the 1980s, they tested more than 400 small molecules in search of one that would block the activity of the errant tyrosine kinase, without derailing other important enzymes. When they found a candidate in 1992, Druker joined the effort and led the way in developing it into Gleevec. **Figure 2** shows how the drug works—it nestles into the pocket on the tyrosine kinase that must bind ATP to stimulate cell division. With ATP binding blocked, cancer cells do not receive the message to divide, and they cease doing so. After passing safety tests, the drug worked so dramatically that it set a new speed record for drug approval—10 weeks.

Erin and the other patients were able to track their progress in several ways:

- "Hematological remission" meant that the percentage of leukemia cells in the blood fell.
- "Cytogenetic remission" meant that the percentage of cells with the Ph¹ chromosome fell.
- "Molecular remission" meant that the level of mRNA representing the fusion gene fell.

Although molecular remission is the goal of CML treatment, in actuality, fusion gene mRNA rarely reaches undetectable levels. As a result, patients can become resistant to Gleevec—relapse occurs in 3 to 16 percent of patients, depending on how sick they were when diagnosed. Resistance is a result of natural selection. Those few cancer cells able to divide in the presence of the drug eventually take over. Again, genetic research came to the rescue. By discovering how

(Continued)

(Concluded)

resistant cells evade the drug, researchers tweaked Gleevec, making it bind more strongly, and developed new drugs that fit the slightly altered active site in resistant cancer cells.

As for Erin, she decided to go off the drug while pregnant. Although she risked relapse, she did not want to expose a fetus to the powerful drug. Her son Alex was born in 2008, healthy. Follow Erin's progress on her blog: <http://www.glamour.com/lifestyle/blogs/editor>.

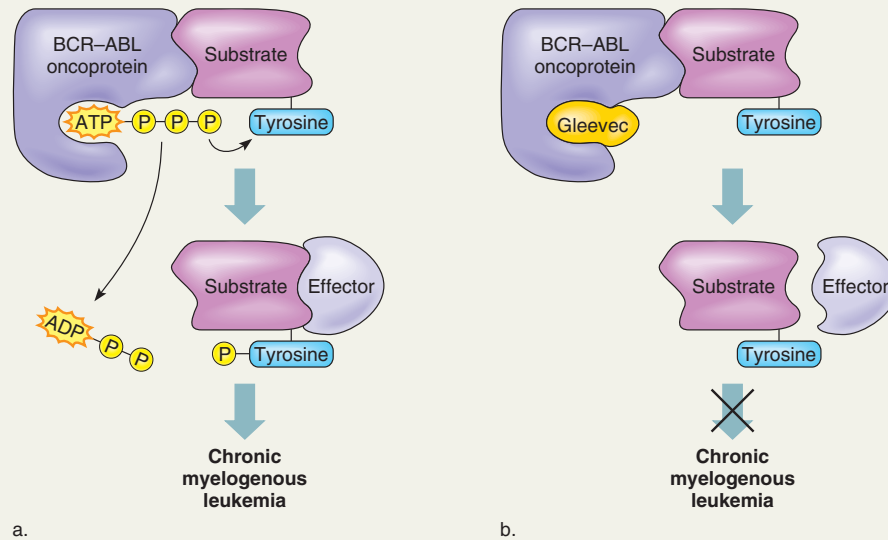


Figure 2 How Gleevec treats chronic myelogenous leukemia. In CML, a translocation forms the fusion oncoprotein BCR-ABL, which functions as a tyrosine kinase. A tyrosine (an amino acid) of a substrate molecule picks up a phosphate from the ATP nestled in the oncoprotein, making the substrate able to bind to another protein, called an effector, that triggers runaway cell division (a). Gleevec replaces the ATP (b). Without phosphorylation of the tyrosine on the substrate, division stops. As cancer progresses, mutations in the DNA of some cells make the shape of their pockets unable to bind the drug. Newer drugs can replace Gleevec once the cancer becomes resistant.

Source: Adapted from "Drug therapy: Imatinib mesylate—A new oral targeted therapy" by Savage & Antman: *New England Journal of Medicine* 346: 683–693. Copyright © 2002 Massachusetts Medical Society. All rights reserved. Reprinted by permission.

p53 Normally Prevents Many Cancers

Another single gene that causes a variety of cancers when mutant is *p53*. Recall from chapter 12 that the *p53* protein transcription factor "decides" whether a cell repairs DNA replication errors or dies by apoptosis. If a cell loses a *p53* gene, or if the gene mutates and malfunctions, a cell with damaged DNA is permitted to divide, and cancer may be the result.

More than half of human cancers involve a point mutation or deletion in the *p53* gene. This may be because *p53* protein is a genetic mediator between environmental insults and development of cancer (figure 18.13). A type of skin cancer, for example, is caused by a *p53* mutation in skin cells damaged by an excessive inflammatory response that can result from repeated sunburns. That is, *p53* may be the link between sun exposure and skin cancer.

In most *p53*-related cancers, mutations occur only in somatic cells. However, in the germline condition Li-Fraumeni syndrome (MIM 151623), family members who inherit a mutation in the *p53* gene have a very high risk of developing cancer—50 percent do

so by age 30, and 90 percent by age 70. A somatic *p53* mutation in the affected tissue results in cancer because a germline mutation in the gene is already present.

Stomach Cancer

Golda Bradfield died of stomach cancer in 1960. By the time some of her grown children developed the cancer too, the grandchildren began to realize that their family had a terrible legacy. Genetic testing revealed that the disease was familial diffuse gastric cancer (MIM 192090). An "exon skipping" mutation deleted DNA from the tumor suppressor gene that encodes E-cadherin, which is essential for cell adhesion in lining cells.

Golda's grandchildren had genetic tests. Eleven of them had not inherited the mutant gene, but scans of their stomachs did not show any tumors. Still, they all had their stomachs removed. It was good that they did, because most of them already had hundreds of tumors, too tiny to have been seen on medical scans. The cousins without stomachs are doing well. Like people who

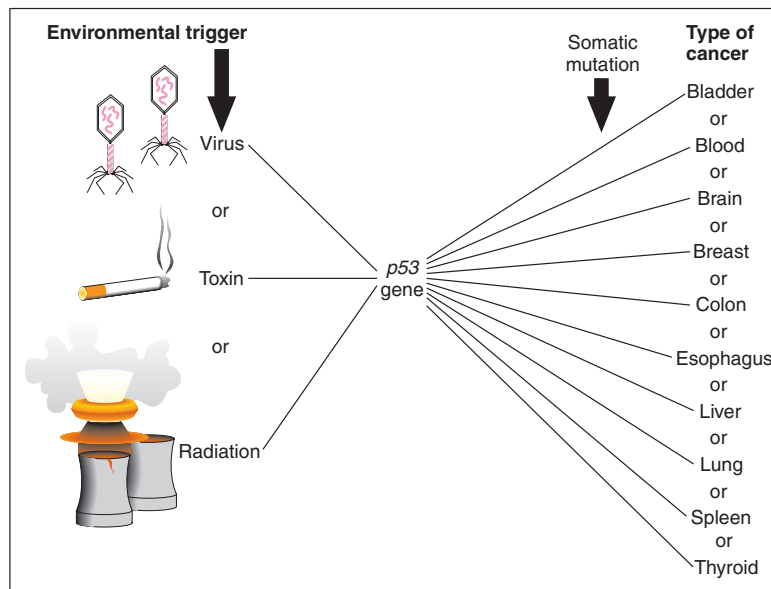


Figure 18.13 *p53* cancers reflect environmental insults. The environment triggers mutations or changes in gene expression that lead to cancer. The *p53* gene is a mediator—“the guardian of the genome.” The protein products of many genes interact with *p53*.

have their stomachs surgically shrunk to lose weight, the cousins have learned to avoid hard-to-digest foods and eat a little at a time, throughout the day. The inconvenience, they say, is a fair trade for eliminating the fear of their grandmother’s, parents’, and aunts’ and uncles’ fates—stomach cancer.

BRCA1

Breast cancer that runs in families may be due to inheriting a germline mutation and then having a somatic mutation occur in a breast cell (a familial form), or two somatic mutations affecting the same breast cell (a sporadic form), as figure 18.4 depicts. However, breast cancer is so common that a family with many affected members may actually have multiple sporadic cases, rather than an inherited form of the disease.

Familial breast cancer exhibits some of the complications of Mendel’s laws described in chapter 5. Mutations in many genes can cause breast cancer, and many are incompletely penetrant, which means that a genotype does not always lead to the associated phenotype. That is, cancer may never develop, even if a person has inherited a breast cancer mutation. For this reason, mutations in breast cancer genes are said to confer susceptibility, rather than cause cancer directly. The genetic heterogeneity of breast cancer also means that absence of one cancer-causing mutation is not a guarantee that breast cancer will never develop.

Only about 5 percent of breast cancers are familial, caused by mutations in any of more than a dozen different genes. Most of the genes associated with familial breast cancer encode proteins that interact in ways that enable DNA to be repaired. If DNA cannot be repaired, mutations that directly cause cancer can accumulate and persist.

The two major breast cancer susceptibility genes are *BRCA1* and *BRCA2*. Together they account for 15 to 20 percent of familial cases. *BRCA1*, which stands for “breast cancer predisposition gene 1,” greatly increases the lifetime risk of inheriting breast and ovarian cancer. This risk, however, varies in different population groups because of the modifying effects of other genes. In the most common *BRCA1* mutation, deletion of two adjacent DNA bases alters the reading frame, shortening the protein. The mutation is inherited as an autosomal dominant trait, with incomplete penetrance and late symptom onset.

BRCA1 and *BRCA2* encode proteins that join two others to form a complex that moves to areas of the genome where both DNA strands are broken at the same site. These double-stranded breaks are particularly dangerous because they cut the chromosome, which means that rearrangements such as deletions and translocation can occur. Normally the protein complex that includes *BRCA1* and *BRCA2* allows *p53* and other cell cycle checkpoint genes to repair the damage or, if it is too extensive, trigger cell death.

BRCA1 mutations have different incidences in different populations. Only 1 in 833 people in the general U.S. population has a mutant *BRCA1* allele. That figure is more than 1 in 50 among Ashkenazi Jewish people, due to population bottlenecks and nonrandom mating. The *BRCA1* gene was initially discovered in Ashkenazi families in which several members developed the cancer at very young ages. In this population group, a woman who inherits a *BRCA1* mutation has up to an 87 percent risk of developing breast cancer over her lifetime, and a 50 percent risk of developing ovarian cancer. Some women in this population group who learn that they have inherited the mutation have their breasts and/or ovaries removed, similar to the cousins who had their stomachs removed to prevent their

family's cancer. However, a person's population group must be considered. The risk of a *BRCA1* mutation causing cancer in the general population is only about 10 percent, based on empirical (observational) evidence. Women with such mutations born after 1940 have a higher risk than those born earlier, suggesting that the environment also plays a role in whether inheriting a mutation causes cancer.

BRCA2 breast cancer is also more common among the Ashkenazim. Ashkenazi women who inherit a mutation in *BRCA2* face a 60 to 85 percent lifetime risk of developing breast cancer and a 10 to 20 percent risk of developing ovarian cancer. Men who inherit a *BRCA2* mutation have a 6 percent lifetime risk of developing breast cancer, which is 100 times the risk for men in the general population. Inheriting a *BRCA2* mutation also increases the risk of developing cancers of the colon, kidney, prostate, pancreas, gallbladder, skin, or stomach.

The fact that *p53*, *BRCA1*, and *BRCA2* proteins all bind to each other in the nucleus suggests that they interact to enable a cell to repair double-stranded DNA breaks. *BRCA2* also pulls apart daughter cells as mitosis completes. Mutations in the *BRCA2* gene may explain the aneuploidy (extra or missing chromosomes) that is common in cancer cells.

Genes whose protein products affect those of *BRCA1*, *BRCA2*, and *p53* can cause breast cancer. For example, the product of a gene called *ATM* adds a phosphate to the product of a gene called *CHEK2*, which then adds a phosphate to the *BRCA1* protein. Mutations in *ATM* and *CHEK2* also cause breast cancer because they affect *BRCA1* protein. Another form of breast cancer results from mutations in any of five genes known to cause Fanconi anemia, a fatal blood disorder. Five of the Fanconi anemia proteins form a cluster that activates a sixth protein, which in turn binds to and inactivates the *BRCA2* protein. Overall, the mutations that increase risk of breast cancer seem to have in common interference with DNA repair.

MicroRNAs Revisited

MicroRNAs are small RNA molecules that act as “dimmer switches,” blocking translation of certain genes into proteins by binding to their mRNA transcripts (see section 11.2). Because microRNAs normally control the expression of certain proto-oncogenes and tumor suppressor genes, when microRNAs themselves have mutations or their expression is too high or too low, cancer can result. For example, in B cell lymphoma, a blood cancer, a B cell has thirty times the normal number of copies of a particular microRNA. Normally, this microRNA regulates activation of B cells in the lymph nodes. The excess copies of the microRNA stimulate too much cell division among the B cells, and cancer results.

The relationship among microRNAs, oncogenes, and tumor suppressor genes is complex and not yet well understood. Cause and effect can be difficult to separate, because oncoproteins and tumor suppressor proteins also affect microRNAs. For example, the tumor suppressor *p53* controls a family of three microRNAs that regulate the cell cycle. Disruption of the function of these microRNAs causes a form of lung cancer. Many of the DNA sequences that encode microRNAs are

found in genome regions that harbor oncogenes and/or tumor suppressors. All of these sequences interact because they target the same pathways.

Changes in microRNA expression also accompany cancer's progression. When cells at the center of a solid tumor become starved for oxygen before angiogenesis brings in an adequate blood supply, a set of microRNAs different from the ones that may have started the cancer appear. These microRNAs alter the expression of yet other genes so that the cell can use metabolic pathways that enable it to survive without oxygen. Yet other combinations of microRNAs alter gene expression as a cancer invades nearby tissue and spreads.

Researchers are using the changes in microRNA expression that accompany cancer to develop new, more sensitive ways to diagnose, treat, and follow response to treatment. For example, different microRNA expression patterns indicate whether pancreatic pain is due to inflammation only, or to cancer—an important distinction. For breast cancer, a microRNA expression “signature” correlates to the likelihood of spread, helping physicians to decide which patients would benefit from further treatments.

At a more fundamental level, understanding microRNAs may solve a mystery in cancer genetics: families in which several members have or had cancer, but do not have any of the recognized, single-gene family cancer syndromes. The fact that a single sequence of microRNA can have many targets—many genes whose translation it blocks—may explain why different family members develop different cancers.

Key Concepts

1. Proto-oncogenes normally control the cell cycle. They can become oncogenes when they mutate, move next to a gene that is highly expressed, or are transcribed and translated with another gene, forming a fusion protein.
2. Mutations in tumor suppressor genes usually are deletions that cause a cell to ignore extracellular constraints on cell division, or affect DNA repair.
3. MicroRNAs alter the expression of oncogenes and tumor suppressor genes. Patterns of microRNA expression change as a cancer progresses.

18.5 Many Genes Contribute to Cancer

The field of cancer genetics has progressed much like genetics in general, from a single-gene approach to considering the sequential and combined actions of many genes.

Genes that guide a cell toward the cancerous state when mutant are considered in two broad categories, based on their effects. “Gatekeeper” genes control mitosis and apoptosis, which must be in balance to maintain the number of cells forming the affected tissue. Their effect is direct. “Caretaker” genes, in contrast, control the mutation rates of gatekeepers,

and may have the overall effect, when mutant, of destabilizing the genome.

Most, if not all, cancers are the culmination of a series of changes in several genes, including gatekeepers and caretakers. To identify the steps and reconstruct the progression of the disease, researchers examine DNA in tumor cells from people in various stages of the same type of cancer. The older the tumor, the more genetic changes have accumulated. A mutation present in all stages acts early in the disease process, whereas a mutation seen only in the tumor cells of sicker people functions late in the process. Each step provides a potential point of treatment.

Colon Cancer

Colon (large intestine) cancer does not usually occur in families with the frequency or pattern of a single-gene disorder. However, when family members with noncancerous growths (polyps) in the colon are considered with those who have colon cancer, a Mendelian pattern emerges. Five percent of colon cancer cases are inherited. One in 5,000 people in the United States has precancerous colon polyps, a condition called familial adenomatous polyposis (FAP; MIM 175100).

Healthy colon lining cells typically live three days. In FAP, they fail to die on schedule and instead build up, forming polyps. FAP begins in early childhood with hundreds of tiny polyps that progress over many years to colon cancer. Both oncogenes and silenced tumor suppressors take part as FAP becomes colon cancer.

Our current understanding of the sequence of genetic changes that underlies FAP colon cancer comes from genealogical sleuthing as well as scientific research on a cluster of families living in Salt Lake City, Utah. Geneticists at the University of Utah compared the vast genealogical records of the Utah Population Database, maintained by the Mormon community, to state registries of cancer cases, births, and deaths. They discovered a founder effect (see section 15.3). The affected families in Utah and another in upstate New York that share the same FAP mutation also share ancestors—specifically, Mr. and Mrs. George Fry. The couple, born in Somerset, England in the 1590s, sailed to Plymouth, Massachusetts with their children around 1630. A son passed the colon cancer mutation to

the branch of the family that settled in upstate New York, and a daughter passed it to the pioneers who left for Utah with other Mormons in the 1850s.

Fast-forward about a century. In the fall of 1947 at the University of Utah, young professor Eldon Gardner told his class that colon cancer might be inherited. A student, Eugene Robertson, excitedly told the class that he knew of a family in which a grandmother, her three children, and three grandchildren had colon cancer.

Intrigued, Gardner delved into the family's records and began interviewing relatives. He eventually found fifty-one family members and arranged for each to be examined with a colonoscope, which is a lit instrument passed into the rectum to view the wall of the colon. The colons of six of the fifty-one people were riddled with polyps, although none of the six had symptoms. Removal of the affected tissue probably saved their lives.

In the years that followed, researchers identified other families with more than one case of colon polyps. Individuals with only polyps were diagnosed with FAP. If a person with colon polyps had cancer elsewhere, extra teeth, and pigment patches in the eye, the condition was called Gardner syndrome, named for the professor.

Researchers identified the chromosomal defect that causes Gardner syndrome in 1985 with the help of a 42-year-old man at the Roswell Park Cancer Institute in Buffalo, New York. He had several problems—no gallbladder, an incomplete liver, an abnormal kidney, mental retardation, and Gardner syndrome. To a geneticist, a seemingly unrelated combination of symptoms suggests a chromosomal abnormality affecting several genes. Sure enough, the man's karyotype revealed a small deletion in the long arm of chromosome 5. This was the first piece to the puzzle of colon cancer. The deletion removed a gene called *APC*. It is the main “gatekeeper” for this type of colon cancer, and is the first step depicted in **figure 18.14**.

Normally *APC* protein binds to another protein, β -catenin, causing a phosphate to be added to it. The phosphorylation prevents β -catenin from acting. But when the *APC* gene is deleted, β -catenin isn't silenced, and instead it enters the nucleus and activates genes that promote mitosis. The cell becomes unable to stop dividing. A tumor forms, but it is not yet malignant. Other pathways, such as those controlled by the

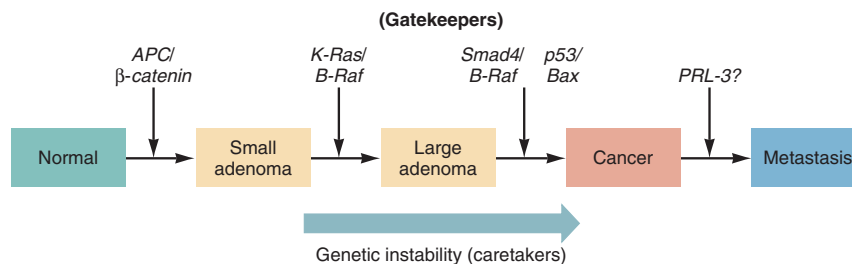


Figure 18.14 Several mutations contribute to FAP colon cancer. Cells lining the colon divide more frequently when the *APC* gene on chromosome 5q undergoes a deletion or point mutation, causing small benign tumors (adenomas) to form. Activation of certain oncogenes, such as *K-Ras* and *B-Raf*, fuel growth of the adenomas. Mutations in *p53* and other genes push the adenoma cells to become cancerous. Finally, mutations in a gene called *PRL-3* trigger metastasis. Caretaker genes cause genetic instability that contributes to the disease process.

genes *TGF* and *p53*, push the abnormal cells to become cancerous. *TGF* normally inhibits mitosis, and *p53* normally sends cells to a fate of apoptosis. *PRL-3* is a gene that acts late in the process, enabling the cancer to spread. Several caretaker genes affect the expression of the gatekeepers, so the overall picture is quite complex.

The Cancer Genome

Several large-scale projects are simultaneously analyzing many aspects of the genomes of cancer cells, constructing descriptive “atlases” that include the types of information in **table 18.3**. A cancer may undergo several dozen genetic changes as it grows and spreads.

In general, many mutations accompany cancer, but they interact in only a few pathways, such as DNA repair in some breast cancers, or cell adhesion in some lung cancers (see table 18.2). Once a pathway is implicated in a particular type of cancer, researchers can look at existing drugs that affect that pathway, or develop new ones that target specific molecules.

A cancer genome study on a form of lung cancer called adenocarcinoma illustrates the type of insights that emerge by cataloging the genetic glitches in a cancer cell. Researchers sequenced 623 genes that affect the pathways involved in cancer in tumor samples from 188 patients, and narrowed the list of mutations detected down to 26 found in a majority of the samples. Most of the 26 implicated genes had not been associated with lung cancer, but had been seen in other types of cancer. These included some already mentioned—*APC* (colon), *ATM* (breast), and *RB* (eye). This study added to the growing

evidence that mutations in the same genes can cause or contribute to cancers in different parts of the body.

Large-scale, genome-level studies of cancer genes enable researchers to investigate environmental effects too. For example, in the adenocarcinoma study, cancer cells from smokers had many more mutations than cancer cells from nonsmokers, and different genes were mutant.

Key Concepts

1. Some cancers may be the culmination of a series of mutations in several genes.
2. Determining which mutations are present in particular stages of a cancer can reveal the sequence of gene actions.
3. Large-scale studies of genetic changes in cancer are revealing shared abnormalities in the same pathways in different cancer types.

18.6 Environmental Causes of Cancer

Environmental factors contribute to cancer by mutating or altering the expression of genes that control the cell cycle, apoptosis, and DNA repair. Inheriting a susceptibility gene places a person farther along a particular road to cancer, but cancer can happen in somatic cells in anyone. Since we cannot do much about our genes, it makes sense to identify environmental cancer triggers and develop ways to control them or limit our exposure to them.

Looking at cancer at a population level reveals the interactions of genes and the environment. For example, researchers examined samples of non-Hodgkin lymphoma tumors for a specific translocation associated with the tumor. The samples were from farmers and had been stored for several years. Of the 172 samples, 65 had the translocation. What did the people who had the translocation have in common that the other stricken farmers didn’t? The individuals who had the translocation were much more likely to have been exposed for long times to toxic insecticides, herbicides, fungicides, and fumigants, compared to the other patients. Therefore, the exposures correlate to the translocation-associated form of the lymphoma.

On an individual level, one way to lower the chance of developing cancer is to avoid certain high-risk environmental factors, such as cigarette smoking and excess sun exposure. A more active approach to minimize environmental influences on cancer risk is chemoprevention, which is taking certain nutrients, plant extracts, or drugs. Promising “chemopreventatives” include folic acid, vitamins D and E, selenium, compounds from soybean, tomato, and green tea, and certain anti-inflammatory drugs. However, a very large study disproved the widely held belief that vitamin E and selenium protect against prostate cancer. The study found that vitamin E in large doses may, in fact, increase the risk. It is important to get information about

Table 18.3 Types of Information in a Cancer “Atlas”

Mutation	<ul style="list-style-type: none"> ■ Somatic or germinal ■ Oncogene or tumor suppressor gene ■ Chromosome level
Variation	<ul style="list-style-type: none"> ■ Single nucleotide polymorphism (SNP) ■ Copy number variant
Gene expression	<ul style="list-style-type: none"> ■ mRNA profile (transcriptomics) ■ Protein profile (proteomics) ■ Epigenetic profile (methylation pattern) ■ MicroRNA pattern
Environmental exposure (carcinogens)	

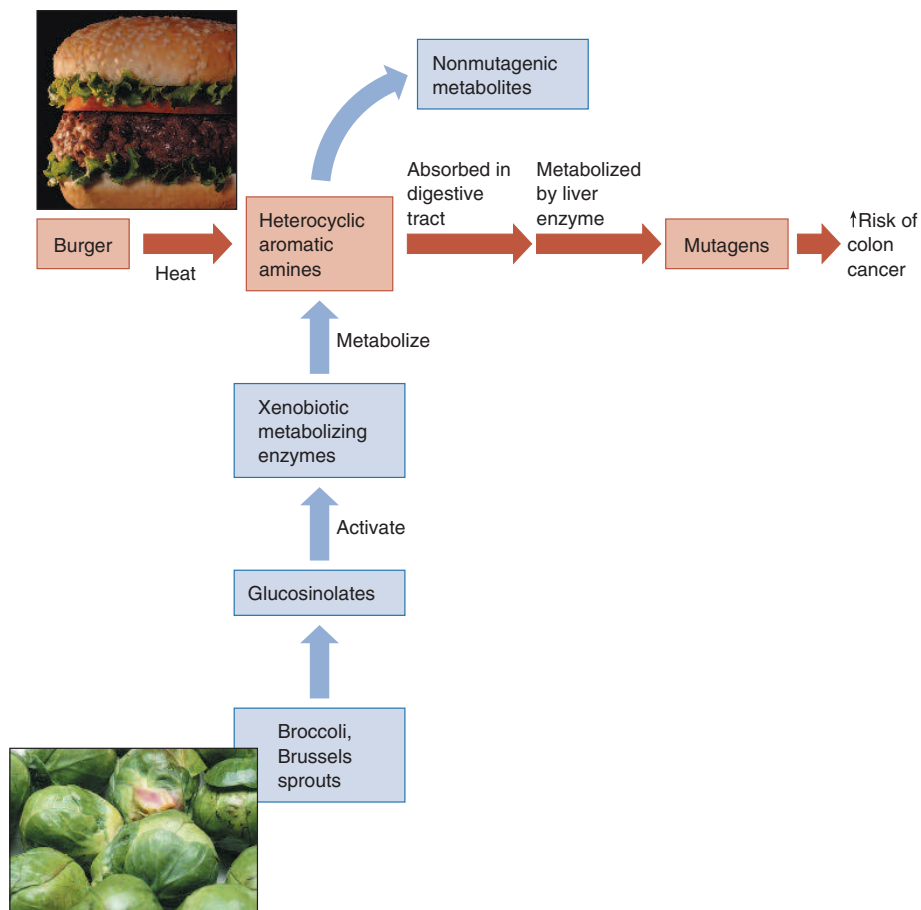


Figure 18.15 One way that cruciferous vegetables lower cancer risk. Compounds called heterocyclic aromatic amines form in cooking meat, are absorbed into the digestive tract, and are metabolized by a liver enzyme into mutagens, which may cause colon cancer. Broccoli and brussels sprouts produce glucosinolates, which activate xenobiotic metabolizing enzymes that block part of the pathway that leads to production of the mutagens.

chemopreventives from health care professionals or medical journals—not from websites selling vitamins.

Considering Carcinogens

Determining precisely how an environmental factor such as diet affects cancer risk can be complicated. Consider the cruciferous vegetables, such as broccoli and brussels sprouts, which are associated with decreased risk of developing colon cancer. These vegetables release compounds called glucosinolates, which in turn activate “xenobiotic metabolizing enzymes” that detoxify carcinogenic products of cooked meat called heterocyclic aromatic amines. With a vegetable-poor, meaty diet, these amines accumulate. They cross the lining of the digestive tract and circulate to the liver, where enzymes metabolize them into compounds that cause the mutations associated with colon cancer (figure 18.15).

Exposure to carcinogens—in the workplace, home, or outdoors—can raise cancer risk. Chemical carcinogens were

recognized as long ago as 1775, when British physician Sir Percival Potts suggested that the high rate of skin cancer in the scrotums of chimney sweeps in London was due to their exposure to a chemical in soot. Since then, epidemiological studies have identified many chemicals as possibly causing cancer in certain populations. However, most studies reveal correlations rather than cause-and-effect relationships. In the strongest cases, genetic or biochemical evidence explains the observed environmental connection.

Methods to Study Cancer-Environment Links

Epidemiologists use different statistical tools to link environmental exposures and cancer. Connections are strengthened when different types of investigations yield consistent results. This is true, for example, of the association between eating whole grain cereals and reduced incidence of colorectal cancer.

Epidemiological studies of cancer causation compare people in different ways. A **population study** compares the

incidence of a type of cancer among very different groups of people. If the incidence differs, then some distinction among the populations may be responsible. For example, an oft-mentioned study from 1922 found that primitive societies have much lower rates of many cancers than more developed societies. The study attributed the lack of cancer to the high level of physical activity among the primitive peoples—but diet might also have explained the difference.

Population studies often have too many variables to clearly establish cause and effect. Consider the very high incidence of breast cancer on Long Island, New York. One hypothesis attributes the mini-epidemic to pesticide exposure, but this population also has a high frequency of *BRCA1* mutations among its Ashkenazi citizens. Sociological factors come into play, too. In this population, women have frequent mammograms starting at a young age. As a result, the percentage of the population with recognized early stages of the disease may be higher than in other populations where women are less likely to have regular mammograms. All of these factors may contribute to the high breast cancer incidence in this area.

More informative than a population study is a **case-control study**, in which people with a type of cancer are matched with healthy individuals for age, sex, and other characteristics. Then researchers look for differences between the pairs. If, for example, the cancer patients had extensive dental X rays at a young age but the control group didn't, X-ray exposure may be a causal factor. Limitations of this type of study are that much of the information is based on recall, people make mistakes, and not all relevant factors are identified and taken into account.

The most informative type of epidemiological investigation is a **prospective study**. Two or more groups of people follow a specified activity plan, such as a dietary regimen, and are checked periodically for cancer. By looking ahead, the investigator has more control over the activities and can verify information. However, a limitation of this type of cancer study is that cancer usually takes many years to appear and progress.

Once epidemiological studies indicate a correlation, a biological explanation is necessary to draw conclusions or suggest further studies. For example, eating certain vegetables that contain antioxidant compounds, which deactivate the free radicals that can damage DNA, may explain the fact that lifetime cancer risk for vegetarians is 29 in 100, compared to 33 in 100 for the general population.

Key Concepts

1. Environmental factors can cause mutations in or alter the expression of genes that raise cancer risk.
2. Avoiding carcinogens and eating foods that detoxify carcinogens can lessen the risks of developing certain cancers.
3. Epidemiologists use population studies, case-control studies, and prospective studies to identify environmental influences on cancer incidence.

18.7 The Personalization of Cancer Diagnosis and Treatment

Estimating the risk that a certain type of cancer will occur in a particular individual is only possible for a few disorders that are inherited in a single-gene fashion through known mutations. More often, discovery of cancer follows a screening test such as mammography or high levels of prostate specific antigen in the bloodstream, or a person feels a lump. Then treatments begin—and there are usually many options.

The oldest cancer treatment is surgery, which prevents invasiveness by removing the tumor. Two other common treatment approaches are radiation and chemotherapy, which kill all cells that divide rapidly. These treatments also affect healthy cells in the digestive tract, hair follicles, and bone marrow, causing side effects of nausea, hair loss, great fatigue, and susceptibility to infection. Other drugs can help patients tolerate the side effects, including colony stimulating factors to replenish bone marrow. These other drugs enable patients to withstand higher and more effective doses of chemotherapy.

Several newer types of cancer drugs affect cancer cell characteristics or activities other than accelerated division rate. Some treatments:

- stimulate cells to regain specialized characteristics, such as drugs based on retinoic acid.
- inhibit telomerase, which prevents cancer cells from elongating their telomeres and continually dividing.
- induce apoptosis, which halts cell division.
- inhibit angiogenesis, which robs a cancer of its blood supply.

The development of angiogenesis inhibitors illustrates the difficulty of pursuing an unusual idea. The term *angiogenesis* was coined by a British surgeon in 1787 to describe the growth of blood vessels in a reindeer's antler, then described in detail in 1935 in the placenta of a pregnant monkey. In 1971, another surgeon, Judah Folkman of Harvard University, suggested that angiogenesis is required to nourish a cancerous tumor, and he tried to block the process. His work was largely unknown to the public until 1998, when a prominent newspaper hyped an off-the-cuff comment that DNA-discoverer James Watson made to a reporter, claiming that Folkman would “cure cancer in 2 years.” He didn't. However, today more than fifty drugs that starve a tumor of its blood supply are available or are being tested. The first angiogenesis inhibitor, Avastin, was approved in 2004 to treat colorectal cancer, in combination with chemotherapy. Angiogenesis inhibitors have several effects: they stop the blood supply to the tumor, they make the tumor soak up more chemotherapy drugs, they kill tumor cells directly, and they stimulate the immune response.

Diagnostic tests and treatments for cancer have become more targeted to cancer cells while sparing healthy ones. The approach to breast cancer illustrates how genetic and genomic information is refining management of these diseases (**table 18.4**). The first targeting came with recognition

Table 18.4 Evolution of Treatments for Breast Cancer

Strategy	Examples
Remove or destroy cancerous tissue	Surgery, radiation, chemotherapy
Use phenotype to select drug	Estrogen receptor-positive women take a selective estrogen receptor modulator or an aromatase inhibitor or both
Use genotype to select drug	Women with <i>HER2</i> -positive cancers take Herceptin (monoclonal antibody)
Genomic level	Gene expression profile on DNA microarray used to guide drug choice; 70-gene signature predicts metastasis

that breast cancer cells have receptors for one or two hormones, estrogen and/or progesterone—or neither receptor type. Women with estrogen receptor-positive tumors begin a several-year course of either a drug that blocks these receptors from receiving signals to divide, or a drug that inhibits an enzyme called aromatase, which is required to produce estrogen.

Determining the estrogen receptor status of a breast tumor is subtyping the disease by phenotype. With the discovery of single genes that cause cancer, diagnosis began to include genotyping. For example, a woman might have *BRCA1* and/or *HER2* breast cancer.

Increasingly, cancer diagnosis utilizes DNA microarrays that scan the genome for cancer-associated mutations as well as detect gene expression patterns. This approach of genetically characterizing tumor cells enables physicians to match a particular patient to the treatments most likely to work right from the start with the fewest side effects, and then to monitor response. DNA tests are also used to predict metastasis. For example, a test that evaluates the expression of seventy genes is used to identify early-stage breast cancers that are most likely to recur after treatment; those tumors are treated more aggressively. The test saves other patients from painful treatment that might not be necessary or effective.

Breast cancer is also described by *lack* of specific receptors. The cancer cells of women with “triple negative” breast cancer do not have excess estrogen, progesterone, or *HER2* receptors. Although these cancers can respond to conventional, nontargeted chemotherapy, the newer drugs that target these receptors are

ineffective. Triple negative breast cancer is aggressive, strikes early, and tends to recur.

The limitation of any cancer treatment, old or new, is defined by the strength of the enemy. Cancer cells are incredibly abundant and ever-changing. Surgery followed by a barrage of drugs and radiation can slow the course of the disease, but all it takes is a few escaped cancer cells—called micrometastases—to sow the seeds of a future tumor. The DNA of cancer cells mutates in ways that enable the cells to pump out any drug sent into them. In addition, cancer cells have redundancies, so that if a drug shuts down angiogenesis or invasiveness, the cell completes the task another way. Although cancer treatments can cure, it is more likely that they kill enough cancer cells, and sufficiently slow the spread, so that it takes

the remainder of a lifetime for the tumors to grow back. In this way, cancer becomes a chronic, manageable condition.

Even as targeted cancer treatments are becoming available, continuing analysis of the human genome is revealing that our view of cancer as a derangement of the cell cycle may be a great oversimplification. Scans of different types of tumors reveal hundreds of genes that mutate as the disease progresses, including well-known cancer genes such as those discussed in this chapter, but also many genes never implicated in cancer before, such as genes that control cell adhesion. In addition, the same type of cancer in different individuals often has different mutations. The overall conclusion: We still have a lot to learn about cancer.

Key Concepts

1. Treatments for cancer target the characteristics of cancer cells. Surgery removes tumors. Chemotherapy and radiation nonselectively destroy rapidly dividing cells.
2. Newer treatments target receptors on cancer cells, block telomerase, stimulate differentiation, or attack a tumor's blood supply.
3. Diagnosis and treatment of cancer will increasingly consider genomic information that enables physicians to better match patient to treatment.

Summary

18.1 Cancer Is Genetic, But Usually Not Inherited

1. Cancer is a genetically dictated loss of cell cycle control, creating a population of highly proliferative cells that invades surrounding tissue.
2. Mutations in genes that encode or control transcription factors, cell cycle checkpoint proteins, growth factors, repair proteins, or telomerase may disrupt the cell cycle, causing cancer. Cancer is polygenic.
3. Changing gene expression patterns also contribute to cancer, and can be used to distinguish cancer types.
4. Sporadic cancers result from two **somatic mutations** in the two copies of a gene. They are more common than cancers that are caused by **germline mutations** plus somatic mutations in affected tissue.

18.2 Characteristics of Cancer Cells

5. A tumor cell divides more frequently or more times than cells surrounding it, has altered surface properties, loses the specializations of the cell type it arose from, and produces daughter cells like itself.
6. A malignant tumor infiltrates tissues and can **metastasize** by attaching to basement membranes and secreting enzymes that penetrate tissues and open a route to the bloodstream. From there, a cancer cell can travel, establishing secondary tumors.

18.3 Origins of Cancer Cells

7. Cell specialization and position within a tissue are important determinants of whether cancer begins and persists.
8. **Cancer stem cells** can divide to yield cancer cells and abnormally differentiated cells.
9. A cell that dedifferentiates and/or turns on expression of “stemness” genes can begin a cancer.
10. A mutation that enables a cell to divide continually can alter the percentages of cells in a tissue that can divide, resulting in an abnormal growth.
11. Chronic repair of tissue damage can provoke stem cells into producing an abnormal growth.

18.4 Cancer Genes and MicroRNAs

12. Cancer is often the result of activation of **proto-oncogenes** to **oncogenes**, and inactivation of **tumor suppressor** genes. Mutations in DNA repair genes cause cancer by increasing the mutation rate.
13. Proto-oncogenes normally promote controlled cell growth, but are overexpressed because of a point mutation, placement next to a highly expressed gene, or transcription

and translation with another gene, producing a **fusion protein**. Oncogenes may also be overexpressed growth factor receptors.

14. A tumor suppressor is a gene that normally enables a cell to respond to factors that limit its division.
15. MicroRNAs control expression of oncogenes and tumor suppressor genes. MicroRNA gene expression changes as a cancer progresses.

18.5 Many Genes Contribute to Cancer

16. To decipher the gene action sequences that result in cancer, researchers examine the mutations in cells from patients at various stages of the same type of cancer. Those mutations present at all stages of the cancer are the first to occur.
17. FAP colon cancer requires several mutations to develop.
18. Cancer atlases catalog all changes in gene structure and function that underlie a particular cancer.

18.6 Environmental Causes of Cancer

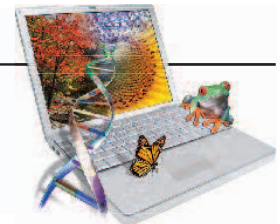
19. **Population, case-control, and prospective studies** can correlate environmental exposures to development of certain cancers. Biochemical and/or genetic evidence can sometimes explain epidemiological observations.

18.7 The Personalization of Cancer Diagnosis and Treatment

20. Traditional cancer treatments are surgery, radiation, and chemotherapy. Newer approaches block hormone receptors, stimulate cell specialization, block telomerase, and inhibit angiogenesis. A genomic approach identifies mutations and differences in gene expression that define cancer subtypes.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

1. Explain the connection between cancer and control of the cell cycle.
2. Explain why not all cells whose chromosomes have long telomeres are cancer cells.
3. Explain how the cell cycle is controlled from both inside and outside the cell.
4. Explain why cancer is usually a genetic disease at the molecular and cellular levels, but not at the whole-body level.
5. Explain why it is important to know whether a cancer is sporadic or inherited.
6. List four characteristics of cancer cells.
7. Define *dedifferentiation*.
8. List the processes and pathways that are abnormal in cancer.
9. Describe what happens as a cancer grows beyond the original tumor.
10. Explain what is inaccurate about the statement “cancer cells are the fastest dividing cells in the body.”
11. Describe four ways that cancer can originate at the cell or tissue level.
12. Define *cancer stem cell*.
13. Distinguish between a proto-oncogene and an oncogene.

14. Describe two ways that oncogenes are activated.
15. Explain how an oncogene is associated with a gain of function and a mutation in a tumor suppressor gene is associated with a loss of function.
16. Explain how retinoblastoma, a *p53*-related cancer, inherited stomach cancer, and *BRCA1* breast cancer have similar causes.
17. Explain how microRNAs can affect oncogenes and tumor suppressor genes.
18. Distinguish between gatekeeper and caretaker genes in cancer.
19. Describe how a form of inherited colon cancer illustrates that several mutations can contribute to causing the disease.
20. Explain how comparing mutations in cells from the same cancer type at different stages can reveal the sequence of genetic changes behind the cancer.
21. Explain how mutations differ from gene expression changes in cancer.
22. Describe a way that exploring the genome of cancer cells reveals information not apparent from focusing on one type of cancer or one gene associated with cancer.
23. Distinguish among population, case-control, and prospective studies to identify environmental factors in cancer.
24. Explain how a cancer treatment that targets cell surface receptors works.
25. Explain why not all cancers affecting the same cell type respond the same way to a particular drug.

Applied Questions

1. An individual can develop breast cancer by inheriting a germline mutation, then undergoing a second mutation in a breast cell; or by undergoing two mutations in a breast cell, one in each copy of a tumor suppressor gene. Cite another type of cancer, discussed in the chapter, that can arise in these two ways.
2. How do the mechanisms of the drugs Gleevec and Avastin differ?
3. A young black woman thinks that she cannot get a *BRCA* form of breast cancer because she isn't Jewish. Is she correct?
4. von Hippel-Lindau syndrome (MIM 193300) is an inherited cancer syndrome. The responsible mutation lifts control over the transcription of certain genes, which, when overexpressed, cause tumors to form in the kidneys, adrenal glands, and blood vessels. Is the von Hippel-Lindau gene an oncogene or a tumor suppressor? Cite a reason for your answer.
5. The *BRCA2* gene causes some cases of Wilms' tumor and some cases of breast cancer. Explain how the same tumor suppressor mutation can cause different cancers.
6. Ads for the cervical cancer vaccine present the fact that a virus can cause cancer as startling news, when in fact this has been known for decades. Explain how a virus might cause cancer.
7. A tumor is removed from a mouse and broken up into cells. Each cell is injected into a different mouse. Although all the mice used in the experiment are genetically identical and raised in the same environment, the animals develop cancers with different rates of metastasis. Some mice die quickly, some linger, and others recover. What do these results indicate about the characteristics of the original tumor cells?
8. Colon, breast, ovarian, and stomach cancers can be prevented by removing the affected organ. Why is this approach not possible for chronic myeloid leukemia?
9. A vegetarian develops pancreatic cancer and wants to sue the nutritionist who suggested she follow a vegetarian diet. Is her complaint justified? Why or why not?
10. MammaPrint is a DNA microarray-based test of the expression of seventy genes implicated in breast cancer. Certain patterns are significantly more common in cancers that spread, creating a "signature" that doctors can use to guide treatment decisions. Cite an advantage and a shortcoming of this test.
11. The discovery of cancer stem cells suggests a new type of treatment—develop a drug that stops self-renewal. Explain how such a drug might work, and what an adverse effect might be.
12. Colorectal cancer is diagnosed in half a million people worldwide each year. In 4 percent of diagnosed individuals, the cancer is part of a familial cancer syndrome, such as Lynch syndrome (MIM 114400). Genetic testing for Lynch syndrome targets mismatch repair genes, and costs about \$3,000. What information would be valuable to decide if it is practical to test for Lynch syndrome for all cases of newly diagnosed colon cancer?
13. A mutation in a gene called *FLT3*, which encodes a tyrosine kinase receptor, causes acute myelogenous leukemia, which has a five-year survival rate of 20 percent. A new drug blocks the receptor on white blood cells. Explain how it works.
14. Rose and Angela are 4-year-old identical twins. Rose develops leukemia, but Angela does not. How is this possible?

Web Activities

15. Go to the Cancer Quest website (www.cancerquest.org). From the menu on the left, click on Cancer Genes. Select an oncogene or tumor suppressor gene and describe how, when mutant, the gene causes cancer.
16. Go to www.cancer.gov. Click on “search for clinical trials.” Select a type of cancer from the drop-down menu, and then click “genetic.” Read about a clinical trial and describe the genetic predisposition or changes that are being studied.
17. Go to <http://cancergenome.nih.gov> and describe a recent discovery. Describe the type of cancer and how genomic information either confirms what was already known about particular genes that cause the cancer, or adds to or changes what was known.
18. Consult the websites for the pharmaceutical companies that market Herceptin, Gleevec, Avastin, or any other cancer drug and explain how the drug works.

Case Studies and Research Results

19. Elsie finds a small lump in her breast and goes to her physician, who takes a medical and family history. She mentions that her father died of brain cancer, a cousin had leukemia, and her older sister was just diagnosed with a tumor of connective tissue. The doctor assures her that the family cancer history doesn't raise the risk that her breast lump is cancerous, because the other cancers were not in the breast. Is the doctor correct?
20. Lung cancer is classified as “small cell” or “non–small cell” based on the appearance of cancer cells under a microscope. However, non–small cell lung cancers fall into three subgroups, based on gene expression patterns. Suggest two ways that this information might be used.

PART 6 Genetic Technology



Transgenic pigs given a bacterial digestive enzyme excrete genetically modified, less-polluting manure. The author poses amidst a pile of the nonmodified material at the University of Georgia.

CHAPTER

19

Genetic Technologies: Amplifying, Modifying, and Monitoring DNA

Improving Pig Manure

Pig manure presents a serious environmental problem. The animals do not have an enzyme that would enable them to extract the mineral nutrient phosphorus from a compound called phytate in grain, so they are given dietary phosphorus supplements. As a result, their manure is full of phosphorus. The element washes into natural waters, contributing to fish kills, oxygen depletion in aquatic ecosystems, algal blooms, and even the greenhouse effect. But biotechnology may have solved the “pig poop” problem.

In the past, pig raisers have tried various approaches to keep their animals healthy and the environment clean. Efforts included feeding animal by-products from which the pigs can extract more phosphorus, and giving supplements of the enzyme phytase, which liberates phosphorus from phytate. But consuming animal by-products can introduce prion diseases, and giving phytase before each meal is costly. A “phytase transgenic pig,” however, is genetically modified to secrete bacterial phytase in its saliva, which enables it to excrete low-phosphorus manure.

A transgenic organism has a genetic change in each of its cells. The transgenic pig has a phytase gene from the bacterium *E. coli*. Its manure has 75 percent less phosphorus than normal pig excrement.

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Knockouts from Gene Targeting

19.1 Patenting DNA

Biotechnology is the use or alteration of cells or biological molecules for specific applications. It is an ancient art as well as a modern science. Using yeast to ferment fruit or produce wine are biotechnologies, as is extracting biochemicals from organisms.

The popular terms “genetic engineering” and “genetic modification” refer broadly to any biotechnology that manipulates DNA. This includes altering the DNA of an organism to suppress or enhance the activities of its own genes, as well as combining the genetic material of different species. Organisms that harbor DNA from other species are termed **transgenic** and their DNA is called **recombinant DNA**.

Creating transgenic organisms is possible because all life uses the same genetic code (**figure 19.1**). Mixing DNA from different species may seem unnatural, but in fact DNA moves and mixes between species in nature—bacteria do it, and it is why we have viral DNA sequences in our chromosomes. But human-directed genetic modification usually endows organisms with traits they would probably not acquire naturally, such as fish that can tolerate very cold water, tomatoes that grow in salt water, and bacteria that synthesize human insulin.

What Is Patentable?

Creating transgenic organisms raises legal questions, because the design of novel combinations of DNA may be considered intellectual property, and therefore patentable. To qualify for patent protection, a transgenic organism, as any other invention, must be new, useful, and not obvious to an expert in the field. A corn plant that manufactures a protein naturally found in green beans but not in corn, thereby making the corn more nutritious, is an example of a patentable transgenic organism. A patent for a

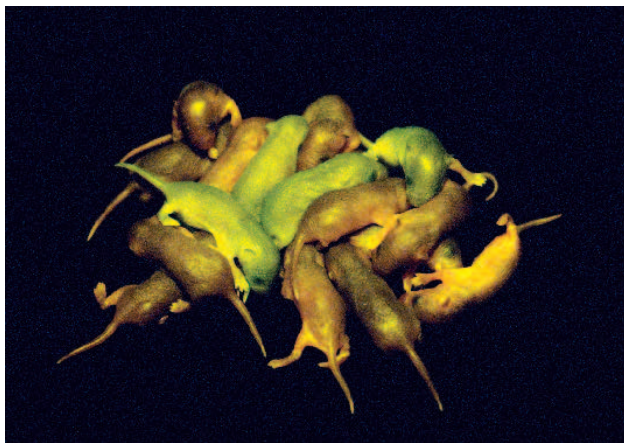


Figure 19.1 The universality of the genetic code makes biotechnology possible. The greenish mice contain the gene encoding a jellyfish’s green fluorescent protein (GFP). Researchers use GFP to mark genes of interest. The GFP mice glow less greenly as they mature and more hair covers the skin. The non-green mice are not genetically modified. The Nobel prize was awarded in 2008 for uses of GFP to mark molecules in live organisms.

DNA sequence from the human genome might be used to diagnose a specific disease, but could inhibit research unless exceptions to use are made for researchers. DNA is also patentable as a research tool, as are algorithms used to extract information from DNA sequences, and databases built of DNA sequences. The Technology Timeline highlights some of the events and controversy surrounding patenting of genetic material.

Patent law has had to evolve to keep up with modern biotechnology. In the 1980s, when sequencing a gene was painstakingly slow, only a few genes were patented. Then, in the mid-1990s, with faster sequencing technology and shortcuts to finding the protein-encoding parts of the genome, the U.S. National Institutes of Health and biotech companies began seeking patent protection for thousands of short DNA sequences, even if their functions weren’t known. Because of the flood of applications, the U.S. Patent and Trademark Office tightened requirements for usefulness. Today, with entire genomes being

Technology Timeline

PATENTING LIFE AND GENES

1790	U.S. patent act is enacted. An invention must be new, useful, and not obvious to earn a patent.
1873	Louis Pasteur is awarded first patent on a life form, for yeast used in industrial processes.
1930	New plant variants can be patented.
1980	First patent is awarded on a genetically modified organism, a bacterium given four plasmids (DNA rings) that enable it to metabolize components of crude oil. The plasmids are naturally occurring, but do not all occur naturally in a single type of bacterium.
1988	First patent is awarded for a transgenic organism, a mouse that manufactures human protein in its milk. Harvard University granted patent for “OncoMouse” transgenic for human cancer.
1992	Biotechnology company is awarded a broad patent covering all forms of transgenic cotton. Groups concerned that this will limit the rights of subsistence farmers contest the patent several times.
1996–1999	Companies patent partial gene sequences and certain disease-causing genes as the basis for developing specific medical tests.
2000	With gene and genome discoveries pouring into the Patent and Trademark Office, requirements tightened for showing utility of a DNA sequence.
2003	Attempts to enforce patents on nonprotein-encoding parts of the human genome anger researchers who support open access to the information.
2007	Patent requirements must embrace new, more complex definition of a gene.
2010	Direct-to-consumer genetic testing companies struggle to license DNA patents for multi-gene and SNP association tests.

sequenced much faster (sometimes in only a matter of days) than it once took to decipher a single gene, a DNA sequence alone does not warrant patent protection. It must be useful as a tool for research or as a novel or improved product, such as a diagnostic test or a drug. In the United States, more than 48,000 patents have been filed on DNA, with 63,000 pending. However, since 1987, only thirty-one lawsuits involving DNA patents have actually gone to court.

Despite the increasing stringency of patent requirements, problems still arise concerning the status of DNA sequences. A biotechnology company in the United States, for example, holds a patent on the *BRCA1* breast cancer gene that includes any diagnostic tests based on the DNA sequence. The American Civil Liberties Union is challenging the *BRCA1* patent, because tests that use the gene are expensive and available only from this one company. The *Bioethics: Choices for the Future* box in chapter 20 explores another patent-related issue: families helping to discover genes by donating their children's DNA, then having to pay for the tests developed from the research.

Analysis of human genome information continues to complicate patenting. One problem is redundancy. For the same gene, it is possible to patent the entire sequence (termed genomic DNA), or just the protein-encoding exons. A researcher can also patent a gene variant, such as a sequence containing a SNP or mutation. A company or researcher developing a tool or test based on a particular gene or its encoded protein might infringe upon several patents that are based on essentially the same information. For example, it is unclear how patents would cover exons common to different genes. Now, as genetics shifts from a gene-by-gene focus to analyzing expression patterns of suites of interacting genes, patent law will have to once again adjust to keep up with scientific developments.

The Patent Thicket

A new problem with patenting DNA stems from the shift in focus of the entire field from a single-gene to a genome-wide approach. A company seeking to use part of a gene sequence in a test must license the use of that sequence from the patent holder. Many companies, however, offer “panels” of tests, such as for several heart-related disorders or for conditions that are more prevalent among Ashkenazi Jews. The companies must license many patented DNA sequences. The challenge is amplified for the many direct-to-consumer genetic testing companies that scan clients' DNA for many thousands of SNPs, looking for “associations” rather than diagnoses. If each SNP is patented, and requires payment of 1 to 5 percent of the profit, such tests cannot be developed unless a company owns the patents.

People in the business of using DNA sequences in tests and products term the need to license patents for every SNP or snippet of DNA the “patent thicket.” Because such companies are forming at a rate that is faster than the Patent and Trademark office can handle the issuing of DNA patents, the U.S.

government has stepped in. A committee has suggested ways around the patent thicket:

- Ban the patenting of associations between DNA sequence variants and disease.
- Allow DNA to be patented only for use in a diagnostic test.
- Exempt physicians and researchers from litigation if they use patented DNA sequences.

A broader action is the Genomic Research and Accessibility Act, which would ban patenting any DNA or its encoded proteins.

While the laws are being worked out, companies can navigate the patent thicket by moving the parts of their operations that use the patented DNA sequences to countries where the restrictions on use do not apply. They can also tweak the recipes for a patented procedure, such as substituting a different type of cell in culture that produces a particular protein, or altering chemical protocols.

The direct-to-consumer companies are finding themselves in an identity crisis. When these companies began to spring up a few years ago, they circumvented regulations on genetic tests for disease by claiming that they provided only information as an educational service. If the law disallows patents for use of DNA sequences in diagnostic tests, these companies would not be included because of how they identify themselves. But if the companies change their tune, claiming to offer tests for diseases so that they have access to patented sequences, they will be under scrutiny of the federal agencies that regulate genetic testing and products for health-related purposes.

Key Concepts

1. Biotechnology is the use or modification of cells or biological molecules for a specific application.
2. DNA patenting is evolving to embrace genome-wide applications.

19.2 Amplifying DNA

Some forensic and medical tests require many copies of a specific DNA sequence from a small sample. Mass-producing a DNA sequence, called nucleic acid amplification, was invented in the 1970s and 1980s. The first and best known technique is the **polymerase chain reaction (PCR)**, which works on DNA molecules outside cells. Another approach, recombinant DNA technology, amplifies DNA from one type of organism placed in the cell of another. Recombinant DNA technology is addressed in section 19.3.

PCR is based on the natural process of DNA replication. Recall from chapter 9 that every time a cell divides, it replicates all of its DNA. PCR uses DNA polymerase to rapidly replicate a specific DNA sequence millions of times.

Applications of PCR are eclectic (**table 19.1**). In forensics, PCR is used routinely to amplify DNA sequences that are profiled to establish blood relationships, to identify remains, and to help convict criminals or exonerate the falsely accused. In agriculture, veterinary medicine, environmental science, and human health

Table 19.1 Uses of PCR**PCR has been used to amplify DNA from:**

- a cremated man, from skin cells left in his electric shaver, to diagnose an inherited disease in his children.
- a preserved quagga (a relative of the zebra) and a marsupial wolf, both extinct.
- microorganisms that cannot be cultured for study.
- the brain of a 7,000-year-old human mummy.
- the digestive tracts of carnivores, to reveal food web interactions.
- roadkills and carcasses washed ashore, to identify locally threatened species.
- products illegally made from endangered species.
- genetically altered bacteria that are released in field tests, to follow their dispersion.
- one cell of an 8-celled human embryo to detect a disease-related genotype.
- poached moose meat in hamburger.
- remains in Jesse James's grave, to make a positive identification.
- the guts of genital crab lice on a rape victim, which matched the DNA of the suspect.
- fur from Snowball, a cat that linked a murder suspect to a crime.

care, PCR is used to amplify the DNA or RNA of pathogens to detectable levels. In genetics, PCR is both a crucial laboratory tool to identify genes and it is a component of many diagnostic tests.

PCR was born in the mind of Kary Mullis on a moonlit night in northern California in 1983. As he drove the hills, Mullis was thinking about the precision of DNA replication, and a way to tap into it popped into his mind. He excitedly explained his idea to his girlfriend and then went home to think it through. "It was difficult for me to sleep with deoxyribonucleic bombs exploding in my brain," he wrote much later.

The idea behind PCR was so simple that Mullis had trouble convincing his superiors at Cetus Corporation that he was onto something. Over the next year, he used the technique to amplify a well-studied gene. Mullis published a landmark 1985 paper and filed patent applications, launching the field of nucleic acid amplification. He received a \$10,000 bonus for his invention, which the company sold to another for \$300 million. Mullis did, however, win a Nobel prize.

PCR rapidly replicates a selected sequence of DNA in a test tube (**figure 19.2**). The requirements include:

1. Knowing parts of a target DNA sequence to be amplified.
2. Two types of lab-made, single-stranded, short pieces of DNA called primers. These are complementary in sequence to opposite ends of the target sequence.
3. A large supply of the four types of DNA nucleotide building blocks.
4. Taq1, a DNA polymerase produced by *Thermus aquaticus*, a microbe that inhabits hot springs. This enzyme is adapted to its host's hot surroundings and makes PCR easy because it does not fall apart when DNA is heated, as most proteins do.

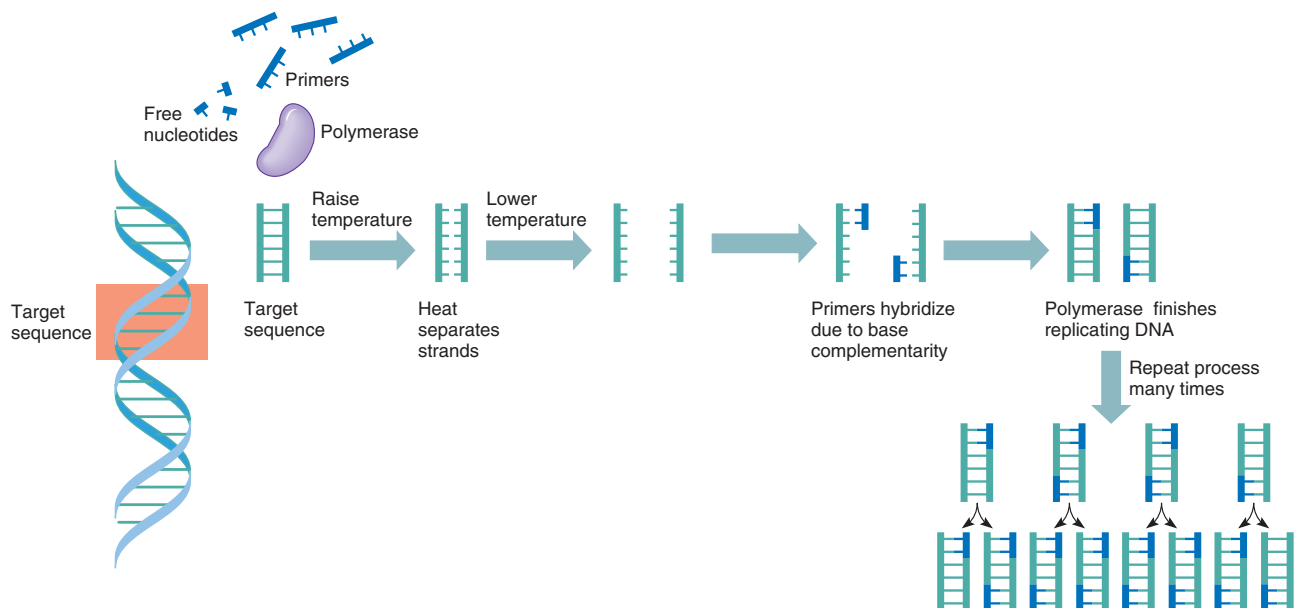


Figure 19.2 Amplifying a specific DNA sequence. In the polymerase chain reaction, specific primers, a thermostable DNA polymerase, and free nucleotides replicate a DNA sequence of interest. The reaction rapidly builds up millions of copies of the target sequence. Figure 14.9 shows an application of PCR.

In the first step of PCR, heat is used to separate the two strands of the target DNA. Next, the two short DNA primers and Taq1 DNA polymerase are added. The temperature is lowered. Primers bind by complementary base pairing to the separated target strands. In the third step, the Taq1 DNA polymerase adds bases to the primers and builds a sequence complementary to the target sequence. The newly synthesized strands then act as templates in the next round of replication, which is initiated immediately by raising the temperature. All of this is done in an automated device called a thermal cycler, or in a device that uses microscopic layers of heated and cooled silicon, to control the key temperature changes. The heat-resistant DNA polymerase is crucial to the process.

The pieces of identical DNA accumulate exponentially. The number of amplified pieces of DNA equals 2^n , where n equals the number of temperature cycles. After just 20 cycles, 1 million copies of the original sequence have accumulated in the test tube.

PCR's greatest strength is that it works on crude samples of rare, old, and minute sequences. PCR's greatest weakness, ironically, is its exquisite sensitivity. A blood sample submitted for diagnosis of an infection, if contaminated by leftover DNA from a previous test, or a stray eyelash from the person running the reaction, can yield a false result.

Using layered silicon instead of a thermal cycler to amplify DNA greatly speeds PCR. Thirty cycles using the thermal cycler take ninety minutes; with the silicon layers, it takes a little over 4 minutes. The speed is valuable in situations where rapid diagnosis is important, such as the case of a person with a life-threatening infection who requires a certain antibiotic, or on a battlefield to detect biological weapons.

The invention of PCR inspired other nucleic acid amplification techniques. One is transcription-mediated amplification, which copies target DNA into RNA and then uses RNA polymerase to amplify the RNA. This procedure doesn't require temperature shifts, and it generates 100 to 1,000 copies per cycle, compared to PCR's doubling, and can yield 10 billion copies of a selected sequence in a half hour.

Key Concepts

1. PCR rapidly replicates a short DNA sequence.
2. PCR is based on DNA replication and has many uses.
3. Other nucleic acid amplification technologies followed PCR.

19.3 Modifying DNA

Recombinant DNA technology adds genes from one type of organism to the genome of another. It was the first gene modification biotechnology, and was initially done in bacteria to produce peptides and proteins useful as drugs. When bacteria bearing recombinant DNA divide, they yield many copies of the "foreign" DNA, and under proper conditions they produce many copies of the protein that the foreign DNA specifies. Recombinant DNA technology is also known as gene cloning.

"Cloning" in this context refers to making many copies of a specific DNA sequence.

Recombinant DNA

Researchers first began pondering the potential uses and risks of mixing DNA from different species in the 1970s. It started in February 1975, when 140 molecular biologists convened at Asilomar, on California's Monterey Peninsula, to discuss the safety and implications of a new type of experiment: combining genes of two species. Would planned experiments that use a cancer-causing virus to deliver DNA be safe? The researchers discussed restricting the types of organisms used in recombinant DNA research and brainstormed ways to prevent escape of a resulting organism from the laboratory. The guidelines drawn up at Asilomar outlined measures of "physical containment," such as using specialized hoods and airflow systems that would keep recombinant microorganisms inside the laboratory, and "biological containment," such as weakening organisms so that they could not survive outside the laboratory.

Recombinant DNA technology turned out to be safer than expected, and it spread to industry faster and in more diverse ways than anyone had imagined. However, recombinant DNA-based products have been slow to reach the marketplace because of the high cost of the research and the long time it takes to develop any new drug. Today, several dozen such drugs are available, and more are in the pipeline. Recombinant DNA research initially focused on providing direct gene products such as peptides and proteins. These included insulin, growth hormone, and clotting factors. However, the technology can target carbohydrates and lipids by affecting the genes that encode enzymes required to synthesize them.

Constructing Recombinant DNA Molecules— An Overview

Manufacturing recombinant DNA molecules requires restriction enzymes that cut donor and recipient DNA at the same sequence; DNA to carry the donor DNA (called cloning vectors); and recipient cells (bacteria or other cultured single cells).

After inserting donor DNA into vectors, the procedure requires several steps to get the desired modified cell type:

- Selecting cells where the genetic material includes any foreign DNA
- Selecting cells that received the gene of interest
- Stimulating transcription of the foreign gene and translation of its protein product
- Collecting and purifying the desired protein

The natural function of restriction enzymes is to protect bacteria by cutting DNA of infecting viruses. Methyl (CH_3) groups shield the bacterium's own DNA from its restriction enzymes. Bacteria have hundreds of types of restriction enzymes. Some of them cut DNA at particular sequences of four, five, or six bases that are symmetrical in a specific way—the recognized sequence reads the same, from the 5' to 3' direction, on both strands of the DNA. For example, the restriction enzyme EcoR1, shown in **figure 19.3**, cuts at the sequence GAATTC. The complementary sequence on the other strand is CTTAAG, which, read

backwards, is GAATTC. (You can try this with other sequences to see that it rarely works this way.) In the English language, this type of symmetry is called a palindrome, referring to a sequence of letters that reads the same in both directions, such as “Madam, I’m Adam.” Unlike the language comparison, however, palindromic sequences in DNA are on complementary strands.

The cutting action of some restriction enzymes on double-stranded DNA creates single-stranded extensions. They are called “sticky ends” because they are complementary to each other, forming hydrogen bonds as their bases pair. Restriction enzymes work as molecular scissors in creating recombinant DNA molecules because they cut at the same sequence in any DNA source. That is, the same sticky ends result from the same restriction enzyme, whether the DNA is from a mockingbird or a maple.

Another natural “tool” used in recombinant DNA technology is a cloning vector. This structure carries DNA from the cells of one species into the cells of another. A vector can be any piece of DNA into which other DNA can insert. A commonly used type of vector is a **plasmid**, which is a small circle of double-stranded DNA that occurs naturally in some bacteria, yeasts, plant cells, and other types of organisms. Viruses that infect bacteria, called bacteriophages, are another type of

vector. Bacteriophages are manipulated to transport DNA but not cause disease. Disabled retroviruses are used as vectors too, as are DNA sequences from bacteria and yeast called artificial chromosomes.

When choosing a cloning vector, size matters. The desired gene must be short enough to insert into the vector. Gene size is typically measured in kilobases (kb), which are thousands of bases. Various types of cloning vectors can hold up to about 2 million DNA bases.

To create a recombinant DNA molecule, a restriction enzyme cuts DNA from a donor cell at sequences known to bracket the gene of interest (**figure 19.4**). The enzyme leaves single-stranded ends on the cut DNA, each bearing a characteristic base sequence. Next, a plasmid is isolated and cut with the same restriction enzyme used to cut the donor DNA. Because the same restriction enzyme cuts both the donor DNA and the plasmid DNA, the same complementary single-stranded base sequences extend from the cut ends of each. When the cut plasmid and the donor DNA are mixed, the single-stranded sticky ends of some plasmids base pair with the sticky ends of the donor DNA. The result is a recombinant DNA molecule, such as a plasmid carrying the human insulin gene. The plasmid and its human gene can now be transferred into a cell, such as a bacterium or a white blood cell.

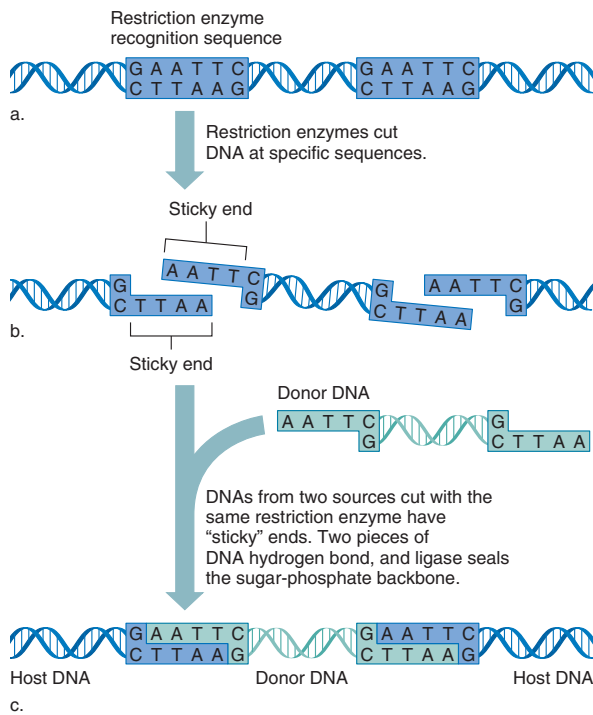


Figure 19.3 Recombining DNA. A restriction enzyme makes “sticky ends” in DNA by cutting it at specific sequences. **(a)** The enzyme EcoR1 cuts the sequence GAATTC between G and A. **(b)** This staggered cutting pattern produces “sticky ends” of sequence AATT. The ends attract through complementary base pairing. **(c)** DNA from two sources is cut with the same restriction enzyme. Pieces join, forming recombinant DNA molecules.

Isolating the Gene of Interest

Constructing recombinant DNA molecules usually begins by cutting all of the DNA in the donor cell. This DNA, which includes non-protein-encoding sequences, is termed genomic DNA. Researchers assemble collections of recombinant bacteria (or other single cells) that harbor pieces of a genome. By using several copies of a genome, the pieces overlap where sequences align. Such a collection is called a **genomic library**. For each application, such as using a human protein as a drug, a particular piece of DNA must be identified and isolated from a genomic library. There are several ways to do this “needle in a haystack” type of search.

A piece of DNA that is complementary to part of the template strand of the gene in question can be linked to a label, such as a radioactive or fluorescent molecule. This labeled gene fragment is called a **DNA probe**. It emits a signal when it binds to its complement in a cell that contains a recombinant plasmid. DNA probes can also be made using genes of similar sequence from other species—they will bind the human version of the gene. Using such a probe is a little like mistakenly typing “hipopotamus” to google “hippopotamus.” You’d probably still come up with a hippo.

A genomic library contains too much information for a researcher seeking a particular protein-encoding gene—it may also contain introns, the genes that encode rRNAs and tRNAs, and many repeated sequences. A shortcut is to use another type of library, called a complementary DNA, or **cDNA library**, that represents only protein-encoding genes. A cDNA library is made from the mRNAs in a differentiated cell, which represent the proteins manufactured there. For example, a muscle cell has abundant mRNAs that encode contractile proteins, whereas a

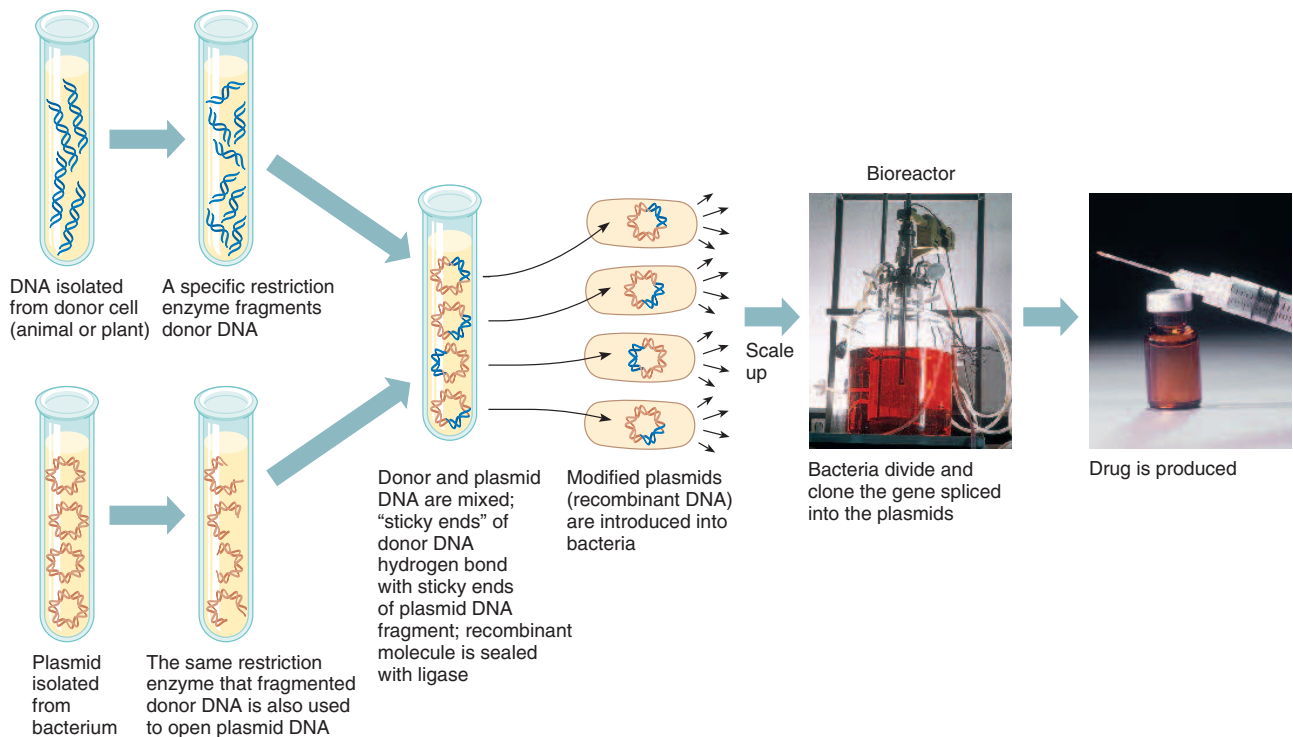


Figure 19.4 Recombinant DNA. DNA isolated from a donor cell and a plasmid are cut with the same restriction enzyme and mixed. Sticky ends from the donor DNA hydrogen bond with sticky ends of the plasmid DNA, forming recombinant DNA molecules. When such a modified plasmid is introduced into a bacterium, it is mass produced as the bacterium divides.

fibroblast has many mRNAs that represent connective tissue proteins.

To make a cDNA library, researchers first extract the mRNAs from cells. Then, these RNAs are used to construct complementary or "c" DNA strands using reverse transcriptase, DNA nucleotide triphosphates, and DNA polymerase (figure 19.5). Reverse transcriptase synthesizes DNA complementary to RNA. DNA polymerase and the nucleotides then can synthesize the complementary strand to the single-stranded cDNA to form a double-stranded DNA. Different cell types yield different cDNA collections, or libraries, that reflect which genes are expressed. They do not, however, reveal protein abundance because in a cell mRNA molecules are transcribed and degraded at different rates.

A specific cDNA can be taken from a cDNA library and used as a probe to isolate the original gene of interest from the genomic library. If the goal is to harness the gene and eventually collect its protein product, then the genomic version is useful, because it includes control regions such as promoters. Once a gene of interest is transferred to a cell where it can be transcribed into mRNA and that RNA can be translated, the protein is collected. Such cells are typically grown in containers called bioreactors, with nutrients sent in and wastes removed. The desired product is collected from the medium in which the cells are growing.

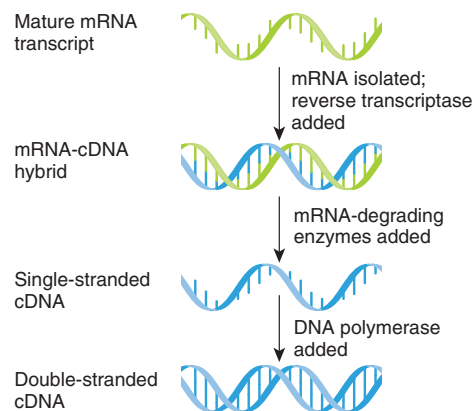


Figure 19.5 Copying DNA from RNA. Researchers make cDNA from mRNA using reverse transcriptase, an enzyme from a retrovirus. A cDNA version of a gene includes the codons for a mature mRNA, but not sequences for promoters and introns. Control sequences from bacteria may be added so that the eukaryotic gene can be transcribed and translated in a prokaryote (the bacterium).

Selecting Recombinant DNA Molecules

Much of the effort in recombinant DNA technology is in identifying and separating cells that contain the gene of interest, once the foreign DNA is inserted into the vector. Three types of recipient cells can result:

1. Cells that lack plasmids
2. Cells that contain plasmids that do not contain a foreign gene
3. Cells that contain plasmids that have picked up a foreign gene (the goal)

The procedure is set up to distinguish bacteria that have taken up recombinant plasmids from those that have not taken up plasmids or that have admitted plasmids that do not carry foreign DNA. One type of strategy has two steps: using an antibiotic resistance gene and a color change reaction to highlight the plasmids that have picked up the gene of interest. First, human and plasmid DNA are cut with the same restriction enzymes and mixed. The plasmids are closed with ligase (the enzyme that glues the sugar-phosphate backbone when DNA replicates), and transferred to bacterial cells. When the antibiotic is applied, only cells harboring plasmids survive. The plasmids also include a gene that encodes an enzyme that catalyzes a reaction that produces a blue color. If a human gene inserts and interrupts the gene for the enzyme, the bacterial colony that grows is not blue, and is therefore easily distinguished from the blue bacterial cells that have not incorporated the human gene.

When cells containing the recombinant plasmid divide, so does the plasmid. Within hours, the original cell gives rise to many cells harboring the recombinant plasmid. The enzymes, ribosomes, energy molecules, and factors necessary for protein synthesis transcribe and translate the plasmid DNA and its foreign gene, producing the desired protein.

Products from Recombinant DNA Technology

In basic research, recombinant DNA technology provides a way to isolate individual genes from complex organisms and observe their functions on the molecular level. Recombinant DNA has many practical uses, too. The first was to mass-produce protein-based drugs.

Drugs manufactured using recombinant DNA technology are pure, and are the human version of the protein. Before recombinant DNA technology was invented, human growth hormone came from cadavers, follicle-stimulating hormone came from the urine of post-menopausal women, and clotting factors were pooled from hundreds or thousands of donors—introducing great risk of infection, especially after HIV and hepatitis C became more widespread.

The first drug manufactured using recombinant DNA technology was insulin. Before 1982, people with type 1 diabetes mellitus obtained the insulin that they injected daily from pancreases removed from cattle in slaughterhouses. Cattle insulin is so similar to the human peptide, different in only two of its fifty-one amino acids, that most people with diabetes could use it. However, about one in twenty patients is allergic to cow insulin because of the slight chemical difference.

Until recombinant DNA technology was developed, the allergic patients had to use expensive combinations of insulin from other animals or human cadavers. **Table 19.2** lists some drugs produced using recombinant DNA technology.

Insulin is a simple peptide and is therefore straightforward to mass-produce in bacteria. Some drugs, however, require that sugars be attached, or must fold in specific, intricate ways to function. These molecules must be produced in eukaryotic cells.

Drugs developed using recombinant DNA technology must compete with conventional products. Deciding whether a recombinant drug is preferable to an existing similar drug is often a matter of economics. For example, interferon β -1b treats a type of multiple sclerosis, but this recombinant drug costs more than \$20,000 per year. British researchers calculated that more people would be served if funds were spent on improved supportive care for many rather than on this costly treatment for a few.

Tissue plasminogen activator (tPA), a recombinant clot-busting drug, also has cheaper alternatives. If injected within 4 hours of a heart attack, tPA dramatically limits damage to the heart muscle by restoring blood flow. It costs \$2,200 a shot. An older drug, streptokinase, is extracted from unaltered bacteria and is nearly as effective, at \$300 per injection. tPA is very valuable for patients who have already had streptokinase and could have an allergic reaction if they were to use it again. *Bioethics: Choices for the Future* on page 387 considers another drug derived from recombinant DNA technology, erythropoietin (EPO).

An application of recombinant DNA technology in the textile industry is a novel source of indigo—the dye used to make blue jeans blue. The dye originally came from mollusks and fermented leaves of the European woad plant or Asian indigo plant. The 1883 discovery of indigo's chemical structure led to the invention of a synthetic process to produce the dye using coal-tar. That method has dominated the industry, but it releases toxic by-products.

In 1983, microbiologists discovered that *E. coli*, with a little help, can produce indigo. The bacterium converts glucose to the amino acid tryptophan, which then forms indole, a precursor to indigo. Another type of bacterium takes the indole to indigo. Researchers altered *E. coli* to suppress alternative pathways for metabolizing glucose, coaxing the cells to synthesize excess tryptophan. They then added genes from the other bacterial species, extending the biochemical pathway all the way to produce indigo. The result: common bacteria that manufacture the blue dye of denim jeans from glucose, a simple sugar.

Transgenic Animals

Eukaryotic cells growing in culture are generally better at producing human proteins than are prokaryotic cells such as bacteria. An even more efficient way to express some recombinant genes is in a body fluid of a transgenic animal, such as milk. The fact that the cells secreting the human protein are part of an animal more closely mimics the environment in the human body.

Table 19.2 Drugs Produced Using Recombinant DNA Technology

Drug	Use
Atrial natriuretic peptide	Dilates blood vessels, promotes urination
Colony stimulating factors	Help restore bone marrow after marrow transplant; restore blood cells following cancer chemotherapy
Deoxyribonuclease (DNase)	Thins secretions in lungs of people with cystic fibrosis
Epidermal growth factor	Accelerates healing of wounds and burns; treats gastric ulcers
Erythropoietin (EPO)	Stimulates production of red blood cells in cancer patients
Factor VIII	Promotes blood clotting in treatment of hemophilia
Glucocerebrosidase	Corrects enzyme deficiency in Gaucher disease
Human growth hormone	Promotes growth of muscle and bone in people with very short stature due to hormone deficiency
Insulin	Allows cells to take up glucose in treatment of type 1 diabetes
Interferons	Treats genital warts, hairy cell leukemia, hepatitis C and B, Kaposi sarcoma, multiple sclerosis
Interleukin-2	Treats kidney cancer recurrence
Lung surfactant protein	Helps lung alveoli to inflate in infants with respiratory distress syndrome
Renin inhibitor	Lowers blood pressure
Somatostatin	Decreases growth in muscle and bone in pituitary giants
Superoxide dismutase	Prevents further damage to heart muscle after heart attack
Tissue plasminogen activator	Dissolves blood clots in treatment of heart attacks, stroke, and pulmonary embolism

Transgenic sheep, cows, and goats have all expressed human genes in their milk, including those that encode clotting factors, clot busters, and collagen. Production of human antibodies in rabbit and cow milk illustrates the potential value of transgenic animals. Recall from figure 17.9 that antibodies are assembled from the products of several genes. Researchers attach the appropriate human antibody genes to promoters for milk proteins. (Recall from chapter 10 that a promoter is a short sequence at the start of a gene that controls the rate of transcription.) These promoters normally oversee production of abundant milk proteins. The mammary gland cells of transgenic animals can assemble antibody parts to secrete the final molecules—just as if they were being produced in an activated B cell in the human immune system. Such antibodies can be used to treat cancer.

Several techniques are used to insert DNA into animal cells to create transgenic animals. They include:

- chemicals that open transient holes in plasma membranes
- liposomes (fatty bubbles) that carry DNA into cells as plasma membranes envelop them
- brief jolts of electricity (electroporation) that open transient holes in plasma membranes
- microscopic needles that inject DNA into cells (microinjection)
- metal particles coated with foreign DNA shot into cells (particle bombardment)

As in plant cells, once foreign DNA is introduced into an animal cell, it must enter the nucleus, replicate along with the cell's own DNA, and be transmitted when the cell divides. Finally, an organism must be regenerated from the altered cell. If the trait is dominant, the transgenic organism must express it in the appropriate tissues at the right time in development. If the trait is recessive, crosses between heterozygotes may be necessary to yield homozygotes that express the trait. Then the organisms must pass the characteristic on to the next generation.

Animal Models

Herds of transgenic farm animals supplying drugs in their milk have not become important sources of pharmaceuticals—they are too difficult to maintain. Transgenic animals are far more useful as models of human disease. Inserting the mutant human beta globin gene that causes sickle cell disease into mice, for example, results in a mouse model of the disorder. Drug candidates can be tested on these animal models and abandoned if they cause significant side effects before testing in humans.

Transgenic animal models, however, have limitations. Researchers cannot control where a transgene inserts in a genome, and how many copies do so. The level of gene expression necessary for a phenotype to emerge may also differ in the model and humans. This was the case for a mouse model of familial Alzheimer disease (MIM 104760). The transgene has the exact same DNA sequence that disrupts amyloid precursor protein in a Swedish family with the condition, but apparently

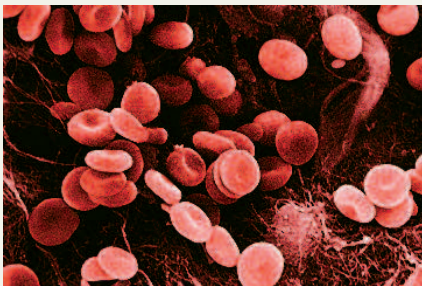
Bioethics: Choices for the Future

EPO: Built-in Blood Cell Booster or Performance-Enhancing Drug?

Athletes are chastised for using performance-enhancing substances. Should this happen if an athlete's body naturally produces greater-than-average amounts of a substance that gives a competitive advantage? This is the situation for EPO (erythropoietin), a glycoprotein hormone that the kidneys produce in response to low levels of oxygen in the blood. EPO travels to the bone marrow and binds receptors on cells that give rise to red blood cell progenitors. Soon, more red blood cells enter the circulation, carrying more oxygen to the tissues (see photo).

The value of EPO as a drug became evident after the invention of hemodialysis to treat kidney failure in 1961. Dialysis removes EPO from the blood, causing severe anemia. But boosting EPO levels proved difficult because levels in human plasma are too low to pool from donors. Instead, in the 1970s, the U.S. government obtained EPO from South American farmers with hookworm infections and Japanese aplastic anemia patients, who secrete abundant EPO into urine. But when the AIDS epidemic came, biochemicals from human body fluids were no longer safe.

Recombinant DNA technology solved the EPO problem. It is sold under various names to treat anemia in dialysis and AIDS



At least two genes control EPO secretion. Certain variants of these genes increase the number of red blood cells, increasing endurance but also raising risk of heart attack and stroke.

patients and is given with cancer chemotherapy to avoid the need for transfusions.

EPO's ability to increase the oxygen-carrying capacity of blood under low oxygen conditions is the reason why athletic training at high altitudes increases endurance. Since the early 1990s, athletes have abused EPO to reproduce this effect, at great risk. EPO thickens the blood, raising the risk of a blockage that can cause a heart attack or stroke, especially when intense, grueling exercise removes water from the bloodstream. Excess EPO caused sudden death during sleep for at least eighteen cyclists abusing the hormone. Olympic athletes are now routinely given urine tests to screen for EPO abuse.

People with familial erythrocytosis get extra EPO naturally. Type 1 (MIM 133100) is autosomal dominant, and is caused by mutation in the EPO receptor. It causes large and abundant red blood cells, but low blood serum levels of EPO. A member of a family from Finland with this condition won several Olympic medals for skiing thanks to his inborn ability. An autosomal recessive form of the condition, type 2 (MIM 263400), increases the level of EPO in the bloodstream. Both forms of erythrocytosis usually have no symptoms, but increase the risk of circulation blocked by the sluggish, oxygen-laden blood.

Questions for Discussion

1. Was it ethical in the 1970s to obtain EPO from sick, poor people in South America and Japan to treat relatively well-off Americans?
2. Should taking a substance made naturally in the body be considered performance enhancement?
3. Do you think that tests should be developed to identify athletes whose genes, anatomy, or physiology give them a competitive advantage in a particular sport or event? What should be done with such information?
4. When developing drugs that use recombinant DNA technology, should researchers consider how the product could be abused?

did nothing to the mice—until researchers increased transcription rate tenfold. Only then did the telltale plaques and tangles, and neuron cell death, appear in the mouse brains.

Animal models might not mimic the human condition exactly because of differences in their rates of development, or different symptoms. Difficulty in relating the abnormal movements and behavior of a mouse model of Huntington disease, for example, led researchers to create transgenic macaques (figure 19.6), which, as primates, are much closer to humans in lifespan, metabolism, reproduction, behavior, and cognition. Recall from chapter 12 that HD is caused by an expansion of a triplet nucleotide repeat. Researchers created macaques with varying numbers of triplet repeats. As in humans, fewer than thirty or so repeats did not affect locomotion or behavior, but longer repeats, or insertion of more than one transgene, produced symptoms.

Sometimes it isn't clear why a transgenic mouse model yields results that do not apply to people who have the exact same mutation. Consider a mouse model of the form of familial amyotrophic lateral sclerosis in which the enzyme SOD1 is not produced (MIM 105400). Ten drugs were found to extend the 2-year lifespan of the mice; none of the drugs helped people, and one actually worsened symptoms. The dog model of ALS in figure 12.1b is not transgenic; it has a dog version of the condition.

Bioremediation

Recombinant DNA technology and transgenic organisms provide processes as well as products. In **bioremediation**, bacteria or plants with the ability to detoxify certain pollutants are released



Figure 19.6 Monkeys stand in for humans. Transgenic macaques are models for Huntington disease, an autosomal dominant neurodegenerative disorder.

or grown in a particular area. Natural selection has sculpted such organisms, perhaps as adaptations that render them unpalatable to predators. Bioremediation uses genes that enable an organism to metabolize a substance that, to another species, is a toxin. The technology uses unaltered organisms, and also transfers “detox” genes to other species so that the protein products can more easily penetrate a polluted area.

Nature offers many organisms with interesting tastes. A type of tree that grows in a tropical rainforest on an island near Australia, for example, accumulates so much nickel from soil that slashing its bark releases a bright green latex ooze. This tree can be used to clean up nickel-contaminated soil.

Bioremediation can tap the metabolisms of transgenic microorganisms, sending them into plants whose roots then distribute the detox proteins in the soil. For example, transgenic yellow poplar trees can thrive in mercury-tainted soil if they have a bacterial gene that encodes an enzyme, mercuric reductase, that converts a highly toxic form of mercury in soil to a less toxic gas. The tree’s leaves then release the gas.

Bioremediation cleans up munitions dumps from wars. One application uses bacteria that normally break down trinitrotoluene—better known as TNT, the major ingredient in dynamite and land mines. The enzyme that provides this capability is linked to the GFP gene. When the bacteria are spread in a contaminated area, they glow near land mines, revealing the locations much more specifically than a metal detector could. Once the land mines are removed, the bacteria die as their food vanishes.

Key Concepts

1. In recombinant DNA technology, a cell receives a cloning vector that contains foreign DNA encoding a protein of interest.
2. Genes are isolated from genomic DNA libraries or cDNA libraries.
3. Antibiotic resistance genes and gene variants that change the color of growth media are used to select cells bearing recombinant plasmids.
4. Recombinant DNA technology is used to manufacture large amounts of a pure protein in single cells and to create multicellular transgenic organisms.
5. Some transgenic plants use *Ti* plasmids to obtain foreign DNA. Transgenic animals receive foreign DNA naked, in liposomes, or by electroporation, microinjection, or particle bombardment. The gene must be transcribed and translated and its product collected and purified. For multicellular organisms, crosses may be necessary to obtain homozygous recessives.
6. Transgenic animals model human disease.
7. Bioremediation uses natural abilities to detoxify environmental contaminants.

19.4 Monitoring Gene Function

Gene expression DNA microarrays (gene chips) are devices that detect and display the mRNAs in a cell. The creativity of the technique lies in choosing the types of cells to interrogate.

Evaluating a spinal cord injury illustrates the basic steps in creating a DNA microarray to assess gene expression. Researchers knew that in the hours after such a devastating injury, immune system cells and inflammatory biochemicals flood the injured area, but it took gene expression profiling to reveal just how fast healing begins.

A microarray is a piece of glass or plastic that is about 1.5 centimeters square—smaller than a postage stamp. Many small pieces of DNA (oligonucleotides) of known sequence are attached to one surface, in a grid pattern. The researcher records the position of each DNA piece in the grid. In many applications, a sample from an abnormal situation (such as disease, injury, or environmental exposure) is compared to a normal control. **Figure 19.7** compares cerebrospinal fluid (CSF; the liquid that bathes the spinal cord) from an injured person (sample A) to fluid from a healthy person (sample B). Messenger RNAs are extracted from the samples

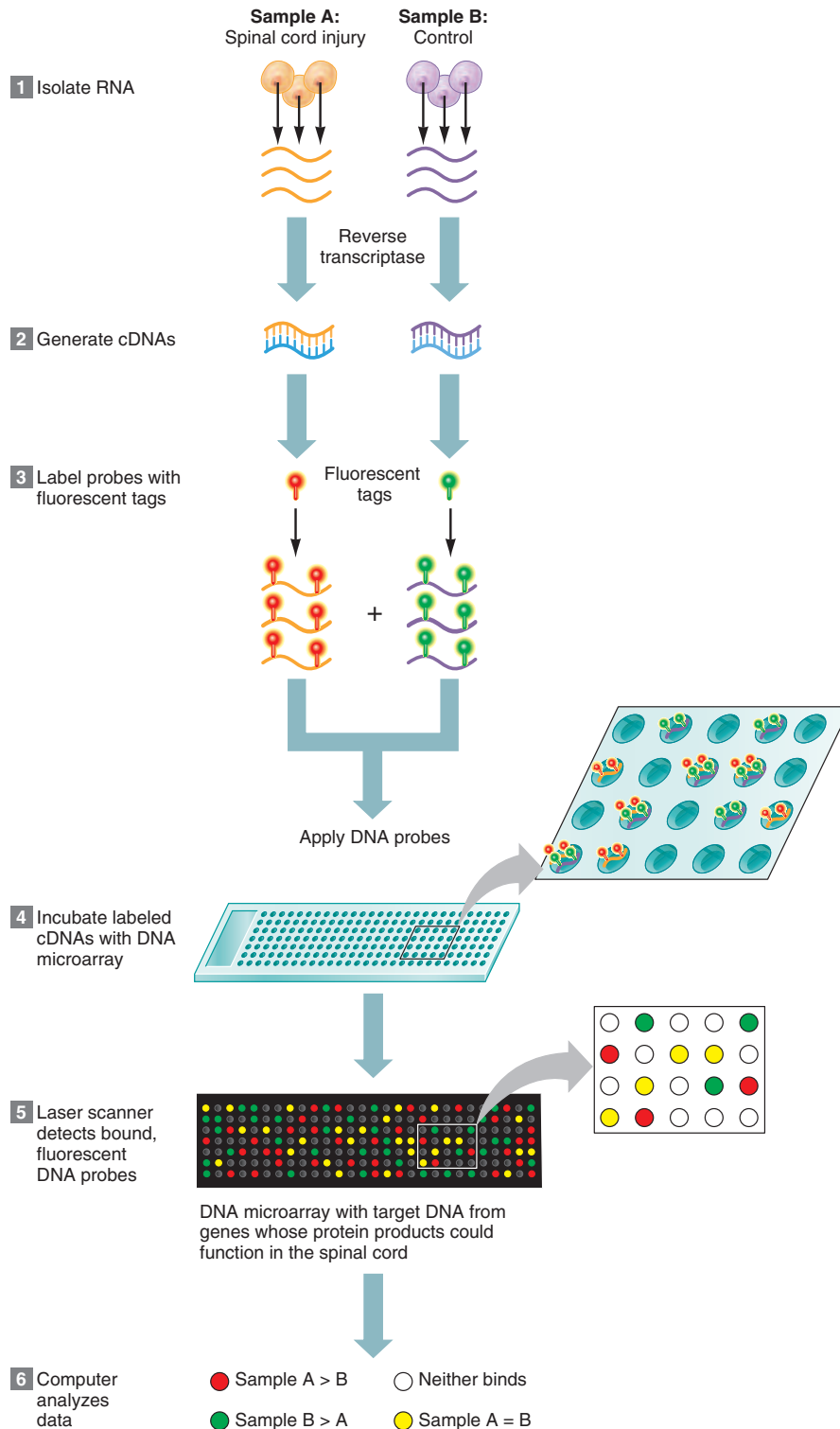


Figure 19.7 A DNA microarray experiment reveals gene expression in response to spinal cord injury.

and cDNAs made (see figure 19.5). The cDNAs from the injury sample are labeled with a red fluorescent dye, and the cDNAs from the control sample are labeled with a green fluorescent dye. These labeled DNAs are then applied to the microarray, which displays thousands of genes likely to be involved in a spinal cord injury, or the entire human genome. This allows for surprises, avoiding the assumption that we know what to look for.

DNA that binds to complementary sequences on the grid fluoresces in place. A laser scanner then detects and converts the results to a colored image. Each position on the microarray can bind DNA pieces from both samples, either, or neither. The scanner also detects fluorescence intensities, which provides information on how strongly the gene is expressed. Then a computer algorithm interprets the pattern of gene expression, which may or may not make visual sense to the researcher—microarray experiments often yield surprise results.

For the spinal cord example, the visual data mean the following:

- Red indicates a gene expressed in CSF only when the spinal cord is injured (and presumably leaking inflammatory molecules).
- Green indicates a gene expressed in CSF only when the spinal cord is intact.
- Yellow indicates positions where both red- and green-bound dyes fluoresce, representing genes that are expressed whether or not the spinal cord has been injured.
- Black, or a lack of fluorescence, corresponds to DNA sequences that are not expressed in CSF.

A computer analyzes the color and intensity pattern, which provides a snapshot of gene expression following spinal cord injury. The technique is even more powerful when repeated at different times after injury. When researchers did exactly that on injured rats, they discovered genes expressed just after the injury whose participation they never suspected. Their microarrays, summarized in **table 19.3**, revealed waves

of expression of genes involved in healing. Analysis on the first day indicated activation of the same suite of genes whose protein products heal injury to the deep layer of skin—a total surprise that suggests new points for drugs to intervene.

Key Concepts

1. DNA microarrays enable researchers to track gene expression.
2. In a DNA microarray experiment, DNA pieces of known sequence are attached to a glass or plastic chip, and differentially labeled sample cDNAs are applied.
3. The patterns and color intensities of spots indicate which genes are expressed. A laser scanner detects and computer algorithm interprets the results.

19.5 Silencing DNA

In some situations, silencing gene expression may be useful, such as blocking transcription of an oncogene or of an enzyme that ripens a fruit, thereby extending shelf life. Chapter 11 (see figures 11.7 and 11.8) discussed microRNAs, which are abundant in cells and are therefore natural ways of controlling gene expression. The techniques described next are introduced means of controlling gene expression.

RNA Interference

The RNA molecules discussed so far—mRNA, tRNA, and rRNA—are single-stranded. However, single-stranded RNA molecules can fold into short, double-stranded regions where the base sequence is complementary, as **figure 19.8** shows. A tRNA molecule in fact assumes its characteristic cloverleaf shape by such localized base pairing. Short, double-stranded RNAs introduced into cells have great effects, and their discovery led to the field of **RNA interference (RNAi)**.

The Nobel Prize in Physiology or Medicine was awarded in 2006 to Andrew Fire and Craig Mello for explaining how RNAi works. They discovered that short, double-stranded RNAs sent into cells separate into single strands, one of which binds its complement in mRNA, preventing it from being translated. This block in translation is RNA interference (RNAi), and the small RNAs that carry it out are called “small interfering RNAs” (siRNAs).

Several proteins and protein complexes orchestrate RNAi, and are also part of microRNA function (**figure 19.9**). First, an enzyme called Dicer cuts long, double-stranded RNAs into pieces twenty to twenty-five nucleotides long. These pieces contact a group of proteins that form “RNA-induced silencing complex,” or RISC, and one strand, called the guide strand, adheres. Now as part of RISC, the guide strand finds its complementary mRNA and binds. Then a protein called argonaute that is part of RISC degrades the targeted mRNA, destroying the instructions for building its encoded protein. SiRNAs act in

Table 19.3 Gene Expression Profiling Chronicles Repair After Spinal Cord Injury

Time After Injury (rats)	Type of Increased Gene Expression
Day 1	Protective genes to preserve remaining tissue
Day 3	Growth, repair, cell division
Day 10	Repair of connective tissues Angiogenesis
Days 30–90	Blood vessels mature New type of connective tissue associated with healing



Figure 19.8 Hairpins. RNA hydrogen bonds with itself, forming hairpin loops.

the nucleus, too, where they alter methylation and the binding of histones to certain genes.

The natural function of RNA interference is not yet fully understood. Perhaps RNA interference is a way that plants and animals combat certain viral infections, which are essentially injections of double-stranded RNA. Even though questions remain, RNAi has spawned a robust biotechnology to “knockdown” gene expression, in which short, synthetic, double-stranded RNAs are used to intentionally destroy particular mRNAs.

RNAi is used to discover gene functions and to silence dangerous gene variants. For example, RNAi-based vaccines knock down expression of key genes in viruses that cause disease, such as AIDS, polio, and hepatitis C. To treat cancer, siRNAs silence gene variants that lift control of the cell cycle. An siRNA can even knockdown an enzyme required for caffeine synthesis in coffee plants, creating a better-tasting decaf.

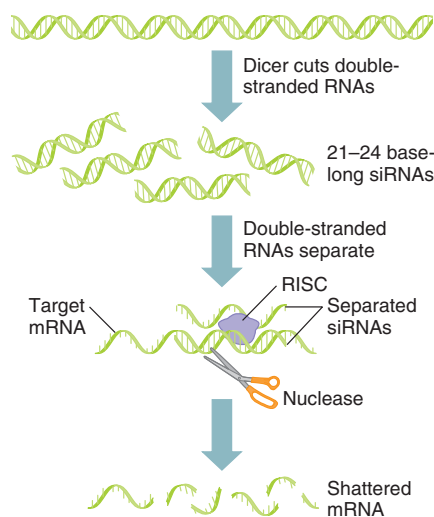


Figure 19.9 RNA interference. Dicer cuts double-stranded portions of RNA molecules, which then associate with RNA-induced silencing complexes (RISCs). The RNAs open, revealing single strands that locate and bind specific mRNAs. Nucleases then break down the targeted mRNAs.

Antisense Sequences

A strategy called “antisense-induced exon skipping” silences mutations that cause exons to be cut out of maturing mRNA (see figure 10.10). This action restores production of the full-length protein. To physically block the mutations from being expressed, researchers introduce synthetic molecules called morpholinos. A morpholino consists of a selected sequence of twenty-five DNA bases that are attached to organic groups that are not the same as the sugar-phosphate backbone of DNA—they are essentially stand-ins for short, complementary sequences that fool the cell by binding to a particular gene part.

Morpholinos are synthesized to be complementary, or “antisense,” to a specific part of a specific gene that includes a mutation that disrupts splicing. They bind to their complements, masking the part of the gene that encodes the part of the mRNA that disrupts exon splicing. In this way, the original mutation is essentially bypassed, enabling some normal protein to be produced.

Silencing mutations with morpholinos has had promising results in treating Duchenne muscular dystrophy (see figure 2.1). Researchers injected morpholinos into the muscles of transgenic mice with DMD, blocking the splicing mutation so that functional dystrophin is made. The treated mice showed improved muscle force. A similar effect was seen in children with DMD when morpholinos were injected into their leg muscles.

Knockouts from Gene Targeting

RNA interference and antisense-induced exon skipping are “knockdown” approaches to silencing gene expression. Knockdown strategies actually evolved from “knockouts,” in which a natural process called homologous recombination is harnessed. It intentionally replaces a DNA sequence with a similar or identical sequence that cannot be transcribed or translated, thereby silencing gene expression. The technique, called gene targeting, was developed in the late 1980s by introducing an inactivated gene into a mouse embryonic stem cell (it does not work on mammalian fertilized ova). Observing what happens (or doesn’t happen) as a mouse embryo develops and the targeted gene does not function can reveal the gene’s normal function as well as present an opportunity to watch the beginnings of an inherited disease. A variation on gene targeting exchanges genes that have an altered function, producing a “knockin.”

Gene targeting to create knockouts has not been used in humans because of the requirement of working with embryos, plus the complex breeding designs that are necessary to get homozygous recessive individuals. However, knockout mice are valuable in several ways:

- They are more accurate models than transgenic mice because a transgene can insert anywhere in a genome. A knockout swaps in a gene at a specific site.
- Populations are easily tested. Two groups of identical knockout mice, for example, can be exposed to different environmental factors to assess their impact.

- Knockouts for several genes can be created to observe polygenic traits and diseases.
- Mouse embryos and fetuses with diseases that humans also get can be observed.

Ironically, the greatest value of knockout mouse models became clear with what looked at first like failed experiments—the knockouts for supposedly vital genes were healthy! For example, knockout mice lacking a collagen gene thought to be essential for normal growth and development of long bones, in both mice and humans, had normal skeletons. These types of results reveal genetic heterogeneity—when different genes have redundant or overlapping effects on the phenotype. Therefore, knocking out one gene's function exposes that of a different gene.

KEY CONCEPTS

1. RNA interference occurs when short, single-stranded RNAs are introduced into a cell and bind to their complements in mRNAs, preventing translation.
2. Antisense-induced exon skipping uses short, DNA-like molecules to block expression of splicing mutations, restoring translation.
3. Gene targeting uses homologous recombination to knock out a specific gene, revealing the gene's function by removing it.

Summary

19.1 Patenting DNA

1. **Biotechnology** alters cells or biochemicals to provide a product. It includes extracting natural products, altering an organism's DNA, and combining DNA from different species.
2. A **transgenic** organism has DNA from a different species. **Recombinant DNA** comes from more than one type of organism. Both are possible because of the universality of the genetic code.
3. Patented DNA must be useful, novel, and non-obvious. Patent law is evolving to ease use of multi-gene tests.

19.2 Amplifying DNA

4. Nucleic acid amplification, such as **PCR**, uses the power and specificity of DNA replication enzymes to selectively mass-produce DNA sequences.
5. In PCR, primers corresponding to a DNA sequence of interest direct polymerization of supplied nucleotides to make many copies.

19.3 Modifying DNA

6. Recombinant DNA technology mass-produces proteins in bacteria or other single cells. Begun hesitantly in 1975, the technology has matured into a valuable method to produce proteins.
7. To construct a recombinant DNA molecule, **restriction enzymes** cut the gene of interest and a **cloning vector** at a short palindromic sequence, creating complementary “sticky ends.” The DNAs are mixed and vectors that pick up foreign DNA selected.

8. **Genomic libraries** consist of recombinant cells containing fragments of a foreign genome. **DNA probes** are used to select genes of interest from genomic libraries. DNA probes may be synthetic, taken from another species, or a **cdNA**, which is reverse transcribed from mRNA.
9. Genes conferring antibiotic resistance and color changes in growth media are used to select cells harboring recombinant DNA. Useful proteins are isolated and purified.
10. A multicellular transgenic organism has an introduced gene in every cell. Heterozygotes for a transgene are bred to yield homozygotes.

19.4 Monitoring Gene Function

11. **DNA microarrays** are devices that hold DNA pieces to which fluorescently labeled DNA probes from samples are applied. They are used in **gene expression profiling**.

19.5 Silencing DNA

12. In **RNA interference**, small interfering RNAs, introduced into a cell, bind their complementary sequences on mRNA, suppressing protein production.
13. Antisense-induced exon skipping blocks expression of splicing mutations, restoring synthesis of a particular protein.
14. Gene targeting uses homologous recombination to knock out a specific gene in a fertilized ovum, revealing gene function.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

- Cite three examples of a DNA sequence that meets requirements for patentability.
- How are PCR and recombinant DNA technology similar, and how do they differ?
- Describe the roles of each of the following tools in a biotechnology:
 - restriction enzymes
 - cloning vectors
 - DNA microarrays
- How are cells containing recombinant DNA selected?
- List the components of an experiment to produce recombinant human insulin in *E. coli* cells.
- Why would recombinant DNA technology be restricted if the genetic code were not universal?
- What is an advantage of a drug produced using recombinant DNA technology compared to one extracted from natural sources?
- Describe three ways to insert DNA into cells.
- Explain the advantages of using a DNA microarray that covers all of the protein-encoding genes in the human genome (the “exome”), rather than selected genes whose protein products are known to take part in the disease process being investigated.
- Compare and contrast the origins and functions of microRNAs and RNA interference.
- Return to chapter 12 and describe the type of mutation that is the basis of antisense-induced exon skipping as a method of gene silencing.
- Compare and contrast the three ways to silence gene expression discussed in section 19.5.

Applied Questions

- Phosphorus in pig excrement pollutes aquatic ecosystems, causing fish kills and algal blooms, and contributes to the greenhouse effect. *E. coli* produces an enzyme that breaks down phosphorus. Describe the steps to create a transgenic pig that secretes the bacterial enzyme, and therefore excretes less polluting feces.
- To diagnose a rare form of encephalitis (brain inflammation), a researcher needs a million copies of a viral gene. She decides to use PCR on a sample of the patient’s cerebrospinal fluid. If one cycle takes 2 minutes, how long will it take to obtain a millionfold amplification?
- HIV infection was once diagnosed by detecting antibodies in a person’s blood or documenting a decline in the number of helper T cells. Why is PCR detection of HIV RNA more sensitive?
- Genetic modification endows organisms with novel abilities. From the following three lists (choose one item from each list), devise an experiment to produce a particular protein, and suggest its use.

Organism	Biological Fluid	Protein Product
pig	milk	human beta globin chains
cow	semen	human collagen
goat	milk	human EPO
chicken	egg white	human tPA
aspen tree	sap	human interferon
silkworm	blood plasma	jellyfish GFP
rabbit	honey	human clotting factor
mouse	saliva	alpha-1-antitrypsin
- Collagen is a connective tissue protein that is used in skincare products, shampoo, desserts, and in artificial skin. For many years it was obtained from the hooves and hides of cows collected from slaughterhouses. Human collagen can be manufactured in transgenic mice. Describe the advantages of the mouse system for obtaining collagen.
- People did not object to the production of human insulin in bacterial cells used to treat diabetes. Yet some people object to mixing DNA from different animal and plant species in agricultural biotechnology. Why do you think that the same general technique is perceived as beneficial in one situation, yet a threat in another?
- A human oncogene called *ras* is inserted into mice, creating transgenic animals that develop a variety of tumors. Why are mouse cells able to transcribe and translate human genes?
- In a DNA microarray experiment, the researcher selects the DNA pieces that are attached to the grid. For example, to study an injury, he or she might choose genes known to be involved in the inflammatory response. How might this approach be limited?
- Devise an experiment using DNA microarrays to determine whether men and women have different hormonal responses to watching an emotional film. (A hormone is a type of messenger molecule that is carried in the blood).

Web Activities

- Use the Web to identify three drugs made using recombinant DNA technology, and list the illnesses they are used to treat.
- Look at the websites for direct-to-consumer genetic testing companies, and discuss the patenting challenges that these companies face in providing tests for multiple genes or SNPs.

12. Go to the DNA Patent Database (<http://dnapatents.georgetown.edu>) and click on "About the DPO." Consider the number of DNA patents filed and granted each year since 2000. Explain why the number of granted patents peaked in 2003 (see chapter 22).
13. Recombinant DNA technology is used to manufacture human growth hormone (hGH), which is used to treat some forms of dwarfism. However, "anti-aging" clinics and websites sell what they claim is hGH to healthy consumers, although studies indicate that the only benefit is a slight increase in muscle mass. Possible side effects are serious, and include diabetes, breast development in men, joint pain, fluid retention, and shortened life. Legislation is pending to classify hGH as a controlled substance, limiting its distribution.
- Consult a website selling hGH and list the claims and warnings. Which do you think are accurate?
 - Do you think laws should restrict access to hGH by people who do not have the medical conditions for which it is prescribed? Cite a reason for your answer.

Case Studies and Research Results

14. Nancy is a transgenic sheep who produces human alpha-1-antitrypsin (AAT) in her milk. This protein, normally found in blood serum, enables the microscopic air sacs in the lungs to inflate. Without it, inherited emphysema results. Donated blood cannot yield enough AAT to help the thousands of people who need it. Describe the steps taken to enable Nancy to secrete human AAT in her milk.
15. To investigate causes of acne, researchers used DNA microarrays that cover the entire human genome. Samples came from facial skin of people with flawless complexions and from people with severe acne. In the following simplified portion of a DNA microarray, one sample is labeled green and comes from healthy skin; a second sample is labeled red and represents skin with acne. Sites on the microarray where both probes bind fluoresce yellow. The genes are indicated by letter and number.





Max Randell is one of the pioneers of gene therapy, which he received directly into his brain to treat Canavan disease. He was not expected to live past age two, yet here he is on his 11th birthday, with his dad, Mike.

CHAPTER

20

Genetic Testing and Treatment

Chapter Contents

- 20.1 Geneticists Find Zebras, and Some Horses**
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- 20.3 Genetic Testing**
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 - DTC Testing and Fuzzy Regulations
 - Nutrigenetics Testing – Buyer Beware
 - Matching Patient to Drug
- 20.4 Treating Genetic Disease**
 - Treating the Phenotype
 - Gene Therapy

Fighting Canavan Disease

Max Randell was not expected to survive his first 2 years. Today, he is on the brink of adolescence, thanks to gene therapy.

In early 1998, Ilyce and Mike Randell, of Buffalo Grove, Illinois, were worried about their 5-month-old son, Max. He could not hold up his head, roll over, reach for objects, or sleep more than an hour, and he was unresponsive. When doctors diagnosed Canavan disease (MIM 271900), which robs brain neurons of their fatty sheaths so that the brain slowly degenerates, they suggested that the best place for the golden-haired boy would be a nursing home. Instead, Max became the youngest person to receive gene therapy for a degenerative brain disease. Today he attends school, gets around in a wheelchair, and communicates by using a computer and an eyetracking system. Although he has to have frequent brain scans and other tests, Max's quality of life has improved—as it has for the dozen other youngsters who followed him in receiving gene therapy for Canavan disease.

Max received his first gene therapy, to test safety, in 1998, and then a more intensive version in 2001. Hundreds of billions of viral particles that each carried a corrected copy of the gene were delivered through six holes drilled into his skull. This particular gene therapy strategy is now being experimentally used for more common brain disorders, such as Parkinson disease. Only about 4,000 children in the world have Canavan disease, (www.canavanresearch.org).

The media often mention gene therapy failures. But Ilyce Randell tells Max's tale whenever a reporter asks. "I cannot even begin to describe the joy we feel just seeing him regain even the slightest bit of functional mobility; he just beams with pride when his body does what he wants it to. None of this would have been possible for Max if he had not received gene therapy."

20.1 Geneticists Find Zebras, and Some Horses

Donna Beales had been sick for almost as long as she could remember. Despite a lack of stamina and frequent illness, she was an active child who loved horseback riding. When her appendix was removed at age 13, X rays revealed malformations of some of her reproductive organs, and only one kidney. At the time, no one connected these anomalies with her other symptoms.

In young adulthood, new symptoms appeared—a nagging cough that didn't respond to standard treatments, impaired vision, and frequent, lingering upper respiratory and digestive infections. For years, Beales saw doctor after doctor, each specialist focusing on one part of her complex condition, none putting the pieces together to paint an accurate diagnosis, because none sought the cause. Finally, an ear, nose and throat specialist put a name on the coughing—angioedema—and suggested that Beales consult a geneticist.

Medical geneticists are trained to see the big picture. They can assemble seemingly unrelated signs and symptoms and the patterns within families, and suggest tests that may identify a gene variant that might be the underlying cause. Dr. Susan Pauker, at a Harvard-affiliated office in Boston, did just that. The birth defects, fatigue, cough combined with recurrent abdominal pain, frequent infections and visual loss, all stemmed from XO (Turner) syndrome and hereditary angioedema type III (HAE III, MIM 610618)—for which Beales had a very rare genotype.

Geneticists lie at the end of many diagnostic journeys, for they practice the opposite of the saying taught to medical students: "*When you hear hoofbeats, think horses, not zebras*" (figure 20.1). This means that the more common explanations for symptoms should be considered first, just as a horse is more common than a zebra. Single-gene disorders, however, are often zebras. Previous chapters have chronicled other searches for a medical explanation that lead to genetics—Hannah Sames and giant axonal neuropathy (*In Their Own Words*, chapter 2) and Bea Reinhoff and her still-unknown condition (chapter 4 Opener). Donna Beales, with her XO syndrome and rare HAE III, calls herself "a zebra among zebras." Her son also has HAE III.



Figure 20.1 Genetic diseases are often zebras. Medical students are told early in their education, "*When you hear hoofbeats, think horses, not zebras.*" However, the most obvious explanation for symptoms—a horse—may not apply to single-gene disorders, which tend to produce signs and symptoms that affect different parts and functions of the body. Often, a geneticist can solve a medical mystery by putting together the pieces.

Genetics professionals are playing an increasingly visible role in discovering and explaining the underpinnings of not only the zebras, but many horses, too. Genetic testing has exploded from obscure tests for rare diseases to tests for common traits and conditions ordered right on one's home computer. This chapter presents the types of genetic services that a health care consumer might encounter: genetic counseling, genetic testing, and gene therapy.

Key Concepts

1. Single-gene disorders often produce a variety of signs and symptoms that can be confusing to health care professionals who are taught to look for the most common causes.
2. Geneticists are increasingly seeking and explaining genetic contributions to common conditions.

20.2 Genetic Counseling

Inherited traits and disorders differ from others in that they may affect more than one person in a family, and with predictable frequencies. The challenges of understanding the diagnosis and repercussions of an inherited condition led to the emergence of the **genetic counselor** (figure 20.2). This health care professional helps individuals and families along the pathway of identifying a genotype that causes or contributes to disease. Genetic counseling addresses medical, psychological, sociological, and ethical issues, and a genetic counselor has medical, scientific, and communication skills.

Reasons to seek genetic counseling:

- Family history of abnormal chromosomes
- Elevated risk of single gene disorder
- Family history of multifactorial disorder
- Family history of cancer



Genetic counseling sessions:

- Family history
- Pedigree construction
- Information provided on specific disorders, modes of inheritance, tests to identify at-risk family members
- Testing arranged, discussion of results
- Links to support groups, appropriate services
- Follow-up contact



Figure 20.2 The genetic counseling process.

A counselor can interpret a DNA test, but also help a patient cope with anxiety, fear, or guilt.

In 1947, geneticist Sheldon Reed coined the term “genetic counseling” for the advice he gave to physician colleagues on how to explain heredity to patients with genetic diseases. In 1971, the first class of specially trained genetic counselors graduated from Sarah Lawrence College, in Bronxville, New York. Today, thirty programs in the United States offer a master’s degree in genetic counselling. In May 1996, the inaugural meeting of the Transnational Alliance for Genetic Counseling united the field, with practitioners sharing challenges unique to their home countries and cultures.

Genetic counseling began in pediatrics and prenatal care, and has become specialized. Counselors today focus on cancer, cardiovascular disease, neurology, or even as distinct a field as ophthalmology or blood disorders. The field has even infiltrated public policy, as genetic testing has become widespread. Genetic counselors in New York, for example, hold “DNA days” to educate state legislators.

Most genetic counselors work directly with patients. This may entail a single visit to explore a test result, such as finding that a pregnant woman is a carrier for cystic fibrosis, or a several-month-long relationship as the counselor guides a decision to take a test for an adult-onset dominant disorder, such as Huntington disease or *BRCA1* breast cancer.

The knowledge that a genetic counselor imparts is similar to what you have read in this book, presented in a personalized manner and applied to a specific disorder. A counselor might explain Mendel’s laws, but substitute a family’s condition rather than discussing pea color inheritance. Or, a counselor might explain how an inherited susceptibility might combine with an environmental factor, such as smoking, to affect health.

A genetic counseling session begins with a discussion of the family’s health history. Using a computer program or pencil and paper, the counselor constructs a pedigree, then deduces and explains the risks of recurrence for particular family members for their particular inherited illness (figure 20.2). Sometimes she will present possibilities and defer discussion of specific risks and options until test results are available. The counselor also explains which second degree relatives— aunts, uncles, and cousins—might benefit from being informed about a diagnosis.

The genetic counselor provides detailed information on the condition and refers the family to support groups. If a couple wants to have a biological child who does not have the illness, a discussion of assisted reproductive technologies (see chapter 21) might be in order.

A large part of the genetic counselor’s job is to determine when specific biochemical, gene, or chromosome tests are appropriate, and to arrange for people to take the tests. The counselor then interprets test results and helps the patient or family choose among medical options. Genetic counselors are also often asked to provide information on drugs that can cause birth defects, although this really is related to development, not genetics. Until recently, people have sought genetic counseling for either of two general reasons: prenatal diagnosis, and a disease in the family.

Prenatal genetic counseling typically presents population (empiric) and family-based risks, explains tests, and discusses whether the benefits of testing outweigh the risks. The couple or woman decide whether amniocentesis, chorionic villus sampling, maternal serum screening, ultrasound, or no testing is appropriate for them. Part of a prenatal genetic counseling session is to explain that tests that rule out some conditions do not guarantee a healthy baby. For example, amniocentesis checks only large-scale chromosome aberrations and single-gene tests must be requested separately. If a test reveals that a fetus has a serious medical condition, the counselor discusses

possible outcomes, treatment plans, and the option of ending the pregnancy.

Counseling when an inherited disease is in a family is another matter. For recessive disorders, the affected individual is usually a child. Illness in the first affected child is often a surprise, and especially if this is a first child, recognition of a problem may take months.

Communicating the risk to subsequent children may be difficult. Many people think that if one child has an autosomal recessive condition, then the next three will be healthy. Actually, each child has a 1 in 4 chance of inheriting the illness. Counseling for subsequent pregnancies requires great sensitivity. Many people will not terminate a pregnancy when the fetus has a condition that already affects their living child, yet some will see that as the kindest option. Genetic counselors must respect these feelings, and tailor the discussion accordingly, while still presenting all the facts.

Counseling for adult-onset disorders does not have the problem of potential parents making life-or-death decisions for existing or future children, but presents the conflicting feelings of people choosing whether or not to find out if a disease is likely in their future. Often, they have seen loved ones suffer with the illness. This is the case for Huntington disease, as the timeline in figure 4.1 and *Bioethics: Choices for the Future* on page 78 depict. Predictive tests are introducing a new type of patient, the “genetically unwell” or those in a “premanifest” state—people with mutant genes but no symptoms (yet). A disease-associated genotype indicates elevated risk, but is not a medical diagnosis, which is based on symptoms.

When genetic counseling began, it was “nondirective,” meaning that the practitioner did not offer an opinion or suggest a course of action, but presented options. That approach is changing as the field moves from analyzing hard-to-treat, rare single-gene disorders to considering inherited susceptibilities to more common illnesses that are more treatable, and for which lifestyle changes might realistically alter outcome. A more recent definition of the role of the genetic counselor is “shared deliberation and decision making between the counselor and the client.”

Providing nondirective genetic counseling can be challenging. One such situation is when a couple has the same autosomal recessive form of deafness, but want to have children, even though they know that the children will be deaf, because they, too, will be homozygous recessive. In one case, when a genetic counselor suggested adoption or intrauterine insemination by donor to avoid deafness, the couple considered the advice a value judgment on their choice to have a child.

In addition to having the counseling skills to help patients, genetic counselors also communicate with physicians and other health care professionals. They are important parts of teams at molecular diagnostic testing laboratories, where they guide physicians in ordering and interpreting tests. Before a test is ordered, the counselor helps to assess the patient’s pedigree, discusses the pros and cons of the appropriate test, and raises ethical issues that might arise when other family members are considered. While the test is underway, the genetic counselor

ensures that time constraints are respected, such as an advancing pregnancy, and updates the physician. Once test results are in, the counselor may request a repeat if they are inconsistent with the patient’s symptoms; interpret the results; suggest additional tests; and alert the physician if the patient might be a candidate for participating in a research project underway or planned at the lab.

The United States has about 3,000 genetic counselors. Because there are so few genetic counselors, and most of them practice in urban areas, access to their services is limited. Finding a genetic counselor with a specific expertise is especially difficult. For example, only 400 genetic counselors in the United States are specially trained in cancer genetic counseling. Due to the shortage of counselors and demand for their services, sometimes other types of health care professionals, such as physicians, nurses, social workers, and PhD geneticists, provide counseling. One survey found that dietitians, physical therapists, psychologists, and speech-language pathologists regularly discuss genetics with their patients.

Some genetic testing companies, both real and Web-based, use “virtual” genetic counseling, in which an interactive computer program calculates risks and provides basic information. Direct-to-consumer genetic testing companies vary widely in their provision of genetic counseling services. Some offer many web pages of information, but without a genetic counselor to interpret the information that a particular test provides. A few companies offer an actual, human genetic counselor available over the phone at any hour.

Other medical professionals may lack or be unaware of the unique combination of skills that a trained genetic counselor offers. For example, a survey of mothers of children with trisomy 21 Down syndrome found many instances of physicians without training in psychology presenting only negative facts to new parents. Said one mother, “The doctor flat out told my husband that this could have been prevented or discontinued at an earlier stage of the pregnancy.” Another woman overheard her doctor call her son an “FLK,” which is medical slang for “funny-looking kid.”

As genetic testing becomes more commonplace, the need for genetic counseling and other genetics-savvy professionals to help individuals and families best use the new information is more compelling than ever.

Key Concepts

1. A genetic counselor provides information to individuals, couples expecting children, and families about modes of inheritance, recurrence risks, genetic tests, and treatments.
2. The counselor helps people make decisions while being sensitive to individual choices.
3. The shortage of genetic counselors has led other health care professionals to provide the service. Genetic counselors help other health care professionals.

20.3 Genetic Testing

Genetic testing is increasingly becoming part of health care, where it is used to diagnose and predict the course of a disease, and predict and monitor response to treatment (table 20.1). Within the next decade it may even become routine to rapidly and inexpensively sequence one's own genome, or to focus on gene variants and markers of clinical importance. *Bioethics: Choices for the Future* in chapter 22 considers uses of affordable personal genome sequencing.

Using genetic tests wisely must balance our tendency to blame all ills on our genes—genetic determinism—with identifying ways to change our lifestyles to compensate for factors that we have inherited and therefore cannot control. The following sections consider types of genetic tests not already discussed.

Newborn Screening

Hunter Kelly was born in 1997, the son of NFL quarterback Jim Kelly and his wife Jill. When he was diagnosed at 9 months of age with Krabbe disease (MIM 245200), also known as galactosylceramide lipidoses after the enzyme that is missing, the damage to his nervous system was already well underway. The enzyme deficiency causes a buildup of certain metabolites, yet too little myelin, the fatty substance that coats neurons. The early symptoms include crying, difficulty feeding, and stiffness. As time goes on, the back arches, the limbs jerk, tube feeding becomes necessary, and motor and mental development slow and stop. The child loses hearing and vision, and usually dies by age 2.

Hunter Kelly lived for 8 years. Had he been born today, Hunter would have been tested for Krabbe disease and dozens of other inborn errors of metabolism, with just a few drops of blood taken from his heel shortly after birth. A cord blood stem



Figure 20.3 Newborn screening expands. Siblings Emma and Matthew Chorey have inherited the same inborn error of metabolism—glutaric acidemia—but Emma was luckier. Thanks to newborn testing, her condition was diagnosed 9 days after her birth, and she was able to begin a special diet to compensate for her metabolic abnormality. This early intervention has prevented buildup of protein in the basal ganglia, the part of the brain that controls movement. She is well. Matthew was diagnosed at 9 months, because newborn testing was not available. He cannot walk or talk but is still an important member of the Chorey family.

cell transplant from a donor could have prevented symptoms. **Figure 20.3** shows a family in which one child suffered, as did Hunter, yet a younger sibling was helped because testing for their disease, glutaric acidemia (MIM 231670), became available before she was born.

Newborn screening tests for inborn errors of metabolism are not genetic tests, but instead they use a technique from analytical chemistry called tandem mass spectrometry to identify

Table 20.1 Types of Genetic Tests

Type of Test	Information Provided	Example
Population carrier screen	Identifies heterozygotes—people with one copy of a mutant gene	College students are offered testing for sickle cell disease carrier status.
Prenatal test	Detects mutant allele in a fetus for a condition present in a family	A couple who know they are carriers of Tay-Sachs disease has a fetus tested.
Prenatal screen	Tests embryos or fetuses from a population for increased risk of a condition, not based on family history	A pregnant woman's blood is tested for elevated level of a protein indicating increased risk for a neural tube defect.
Newborn screen	Populationwide testing for several treatable inborn errors of metabolism	A child with identified sickle cell disease genes at birth can ease or delay symptoms with antibiotics.
Diagnostic test	Confirms diagnosis based on symptoms	A child with "failure to thrive" and frequent lung infections is tested for mutant alleles for CF.
Predisposition test	Detects allele(s) associated with an illness, but not absolutely diagnostic of it	A young Jewish woman with a strong family history of breast cancer has a mutant <i>BRCA1</i> allele, giving her an 86 percent lifetime risk of developing the condition.
Predictive test	Detects highly penetrant mutation with adult onset in an individual at high risk based on family history	A healthy person is tested for the Huntington disease mutation because a parent has the condition.

unusual metabolites or chemical imbalances that indicate a particular disease. “Mass spec” can detect telltale biochemicals from many disorders at a time, represented in blood drops from a newborn’s heel. In the United States, the individual states determine which diseases to test for. The American College of Medical Genetics recommends that a minimum of twenty-nine disorders be tested for in newborns, but the actual number of tests ranges from twenty-one to fifty-four. This spotty regulation means that a child in one state may be diagnosed in time for treatment, but not those in a different state. Even testing for disorders that currently lack treatment is useful, because it can help narrow down a diagnosis.

The field of newborn screening began in 1961, with phenylketonuria, discussed in section 5.2. The Guthrie test sampled blood from a newborn’s heel and tested for the amino acid buildup that indicates PKU. In 1963, a specialized diet (legally termed a “medical food” so that insurance will cover the high cost) became available, with dramatic positive results. The diet sharply reduces the amount of the amino acid phenylalanine, which builds up in the disorder. The diet is very difficult to follow, and must be for many years, but it does prevent mental retardation. After the success of newborn screening for PKU, state testing expanded to include eight genetic conditions and a nongenetic form of hearing loss. Gradually, the offerings have grown.

Newborn screening is important because the earliest signs and symptoms of many of the disorders tested for are common in babies, and an inexperienced parent may be told “oh, it’s just colic,” and sent home, only to witness declines in motor and mental skills that indicate a problem. Often, treatment must begin in the first weeks after birth to help. Another early symptom for many inborn errors of metabolism is a strange scent to the child’s urine. This is, in fact, what led an astute mother in Norway to mention to a biochemist friend that her baby’s urine smelled “mousy”—a sign of PKU that eventually led to understanding the biochemical basis of the disease. “Maple syrup urine disease” has an obvious odor. Other inborn errors cause the urine to smell like sweaty feet, boiled cabbage, chlorine, or rotting fish.

Some states perform DNA tests on newborns as well as biochemical tests. The National Newborn Screening and Genetics Resource Center helps states to tailor programs to specific populations. For example, a test for cystic fibrosis might include 100 mutations out of the 1,600⁺ that have been identified, but the 100 most common mutations in New Mexico differ from those in New Jersey.

DTC Testing and Fuzzy Regulations

Companies market DNA-based tests for traits, susceptibilities, and genetic diseases, to the general public (**table 20.2**). The tests range from the obvious (eye color) to the dubious (athletic ability) to the serious (cancers).

In the United States, the Clinical Laboratory Improvement Amendments, or CLIA, control genetic testing of body materials for the prevention, diagnosis, or monitoring response to treatment of a disease or health impairment. The CLIA

regulations, instituted in 1988, added “specialty areas” in 1992 to cover very complex tests, such as those involving immunology or toxicology. These did not include genetic testing, which was very limited. Discoveries of the genes behind some of the more common single-gene disorders were just beginning. Since then, the genetics community has repeatedly asked that genetic tests be included as a specialty area, to no avail. State regulations can override CLIA, but only if they are equally or more stringent. Therefore, the regulation of genetic tests remains somewhat unclear. Adding to the confusion is that few people, in the mid-1990s, could have imagined that consumers would be able to order DNA tests on home computers.

Tests that offer genetic information, and not diagnoses, do not come under the CLIA regulations. The distinction between information and diagnosis is often based on careful wording. Consider “direct-to-consumer” (DTC) Company B in table 20.2, which “. . . scans your DNA for genetic risk markers associated with both common and uncommon health conditions.” “Association” means a relationship between one piece of information and another—it is not a correlation or a cause, nor a diagnosis. Some companies have CLIA certification for some of their tests but not others, but even that regulation is not always clear. For example, CLIA certifies that laboratories are safe and that a particular test measures what it claims to measure. However, the regulations do not require that knowing the measurement, such as the concentration of a molecule in blood, leads to actions that improve health.

Even DTC tests for very well-studied mutations can lead to complications and confusion. This is the case for hereditary hemochromatosis (HH, MIM 235200). In this autosomal recessive “iron overload” disease, cells in the small intestine absorb too much iron from food. Early signs and symptoms of HH include chronic fatigue, increased susceptibility to infection, hair loss, infertility, muscle pain, and feeling cold. Over many

Table 20.2 Direct-to-Consumer “Genetic Association” Tests

DTC website/companies that offer information may not be held to the same standards as laboratories that provide tests used to make clinical diagnoses. Here are some offerings from two DTC companies.

Company A - Traits	Company B - Disorders
Alcohol flush reaction	Alzheimer disease
Athletic performance	Brain aneurysm
Bitter taste	Cancers
Earwax type	Diabetes
Eye color	Glaucoma
Lactose intolerance	Multiple sclerosis
Muscle performance	Obesity
Skin quality	Osteoarthritis, restless legs syndrome, rheumatoid arthritis

years, the excess iron is deposited throughout the body, damaging vital organs. However, HH is incompletely penetrant—that is, many people who have the disease-associated genotype do not have symptoms, especially women who lose the extra iron in the monthly menstrual flow.

Diagnosis of symptomatic HH is important, because lowering the body's iron levels is easy—have blood removed periodically. However, diagnosis is based on an increase in the level of the iron-carrying protein ferritin in the blood and confirmed with a liver biopsy, rather than on a genetic test because of the incomplete penetrance. But a website offering genetic testing says: “Most patients do not know that they can test themselves for hereditary hemochromatosis. Here is an easy way to get direct testing without a prescription and results sent directly to the patient only.” At-home testing for HH could be confusing to consumers unfamiliar with the uncertainty of genetics.

Nutrigenetics Testing—Buyer Beware

Some direct-to-consumer testing company websites offer genetic tests along with general questionnaires about diet, exercise, and lifestyle habits. The company then sends a “nutrigenetics” profile with dietary suggestions—often with a pitch to purchase a pricy package of exactly the supplements that an individual supposedly needs to prevent realization of his or her genetic fate.

After the media spread the word of these services, the U.S. Government Accountability Office tested the tests. An investigator took two DNA samples—one from a 9-month-old girl and the other from a 48-year-old man—and created fourteen lifestyle/dietary profiles for these “fictitious consumers”—twelve for the female, two for the male. The samples were sent to four companies that advertise nutrigenetics testing on the web. Here is an example of the information sent to the companies:

- The DNA from the man was submitted as being from a 32-year-old male, 150 pounds, 5'9", who smokes, rarely exercises, drinks coffee, and takes vitamin supplements.
- The DNA from the baby girl was submitted as being from a 33-year-old woman, 185 pounds, 5' 5", who smokes, drinks a lot of coffee, doesn't exercise, and eats a lot of dairy, grains, and fats.
- The same baby girl DNA was also submitted as that of a 59-year-old man, 140 pounds, 5' 7", who exercises, never smoked, takes vitamins, hates coffee, and eats a lot of protein and fried foods.

The elevated risks found for the three were exactly the same: osteoporosis, hypertension, type 2 diabetes, and heart disease. One company would provide the appropriate multi-vitamin supplements for \$1,200, which the investigation found to be worth about \$35. Recommendations tended to state the obvious, such as advising a smoker to quit. The advice tracked with the fictional lifestyle/diet information, and not genetics. Concluded the study: “Although these recommendations may be beneficial to consumers in that they constitute common sense health and dietary guidance, DNA analysis is not needed

to generate this advice.” Some of the suggestions could even be dangerous, such as vitamin excesses in people with certain medical conditions. (A health history was not required.)

Matching Patient to Drug

People react differently to the same dose of the same drug because we differ in the rates at which our bodies react to drugs and break them down. Genetic tests can highlight these differences. Specifically, a **pharmacogenetic** test detects a variant of a single gene that affects drug metabolism, and a **pharmacogenomic** test detects variants of multiple genes or gene expression patterns that affect drug metabolism. Pharmaceutical and biotechnology companies now routinely use these tests in developing drugs, and physicians are increasingly using these tests in prescribing drugs. Pharmacogenetics and pharmacogenomics are often considered together under the umbrella term “personalized medicine.”

Genetic testing to guide drug selection offers several advantages:

- Identifying patients likely to suffer an adverse reaction to a drug.
- Selecting the drug most likely to be effective.
- Monitoring response to drug treatment.
- Predicting the course of the illness (prognosis).

One of the first applications of pharmacogenetics was in breast cancer (see table 18.4), where women with the *HER2* subtype respond to the drug Herceptin. A pharmacogenomic example is the use of a DNA microarray depicting the expression of eighteen genes to predict whether a person is likely to respond to the drugs used to treat hepatitis C, which have severe side effects. A pharmacogenetic/genomic approach might have averted disaster in 2004, when widespread use of an arthritis medication called a COX-2 inhibitor caused heart damage in some patients. Several drugs were discontinued or their use restricted, robbing many people with arthritis of their benefits. Tests that detect specific variants of genes that encode proteins called cytochromes (P450, 2D6, and 2C19) are now used to predict who will develop adverse effects from these drugs, so that prescribing can be appropriately restricted.

One of the first widespread uses of pharmacogenetics is in prescribing the blood thinner Warfarin (also known as coumadin). This drug has a very small range of concentration in which it keeps blood at a healthy consistency, but people can vary up to tenfold in the dose required. Too little allows dangerous clotting; too much causes dangerous bleeding. In the past, physicians would give an initial standard dose, based on a patient's age, gender, health status, weight, and ethnicity, then monitor the patient for a few weeks to check for too much clotting or bleeding, tweaking the dose until it was about right. But this general approach led to hospitalization for abnormal bleeding in 43,000 of the 2 million people prescribed the drug each year.

A “pharmacogenetic algorithm” is now used to prescribe Warfarin. It considers two genes: two variants of *CYP2C9* and one variant of *VKORC1* are associated with

increased sensitivity to the drug. People with these gene variants require lower doses of Warfarin. The new, genetic way of testing for Warfarin response is especially helpful for the 50 percent of patients who fall at the extremes of the range of drug concentration that is effective.

Key Concepts

1. Newborn screening uses tandem mass spectrometry to detect dozens of telltale metabolites in a single blood sample. Newborns undergo DNA testing too. States vary in the number of newborn tests they require.
2. Government regulations for genetic testing apply to labs that produce DNA-based tests used to diagnose disease, but do not apply to tests that provide information on associations. Unawareness of incomplete penetrance is one complication of DTC genetic testing.
3. Nutrigenetics testing can be very misleading.
4. Pharmacogenetics and pharmacogenomics help physicians to select the best drugs for individual patients.

20.4 Treating Genetic Disease

The number of tests for genetic diseases is much greater than the number of treatments. A great challenge with developing such treatments is ensuring that they correct the abnormality in the appropriate cells and tissues to prevent or minimize symptoms. Treatments have evolved through several stages, in parallel to development of new technologies:

- Removing an affected body part
- Replacing an affected body part or biochemical with material from a donor
- Delivering pure, human proteins derived from recombinant DNA technology to compensate for the effects of a mutation
- Gene therapy, to replace mutant alleles

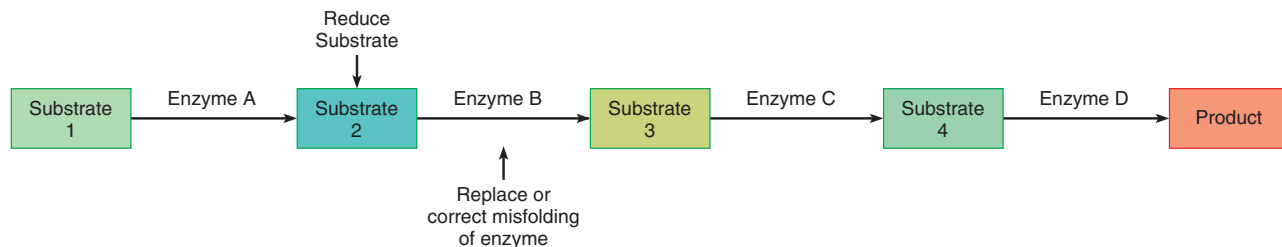


Figure 20.4 Counteracting a metabolic abnormality. Treatments of lysosomal storage diseases are based on understanding metabolic pathways, in which a series of enzyme-catalyzed reactions leads to formation of a product. If any enzyme is deficient or its activity blocked, the substrate builds up and the product is deficient. Enzyme replacement therapy delivers an absent or deficient enzyme. Substrate reduction therapy decreases the amount of substrate and pharmacological chaperone therapy corrects misfolded proteins.

The first three approaches affect the phenotype. Only gene therapy attempts to alter the genotype.

Treating the Phenotype

Preventing a disease-associated phenotype may be as simple as a child with cystic fibrosis adding to her applesauce the digestive enzymes that her pancreas cannot make, or a boy with hemophilia taking a clotting factor to prevent bleeding. Inborn errors of metabolism are particularly treatable when the biochemical pathways are well understood and enzymes can be replaced.

Lysosomal storage diseases are a subclass of inborn errors of metabolism whose effects on cells are well known—a deficient or abnormal enzyme leads to buildup of the substrate (the molecule that the enzyme acts on) as well as a deficit of the breakdown product of the substrate. Recall from figure 2.6 that a lysosome is an organelle that functions as a garbage can of sorts in a cell. It houses forty types of enzymes, and each breaks down a specific molecule.

Type 1 Gaucher disease (MIM 230800) is a lysosomal storage disease that illustrates three general approaches to counteracting an inborn error of metabolism that affects an enzyme, summarized in **table 20.3** and **figure 20.4**. In type 1 Gaucher disease, the enzyme glucocerebrosidase is deficient or absent. As the substrate builds up because there is little or no enzyme to break it down, lysosomes swell, ultimately bursting

Table 20.3

Treatments for Lysosomal Storage Diseases

Treatment	Mechanism
Enzyme replacement therapy	Recombinant human enzyme infused to compensate for deficient or absent enzyme
Substrate reduction therapy	Oral drug that reduces level of substrate so enzyme can function more effectively
Pharmacological chaperone therapy	Oral drug that binds to patient's misfolded protein, restoring function

cells. Symptoms include an enlarged liver and spleen, bone pain, and deficiencies of blood cells. Too few red blood cells cause the fatigue of anemia; too few platelets cause easy bruising and bleeding; and too few white blood cells increase the risk of infection. The disease is very variable in age of onset, severity of symptoms, and rate of progression.

Early treatments for Gaucher disease corrected affected body parts: removing the spleen, replacing joints, transfusing blood, or transplanting bone marrow. In 1991, *enzyme replacement therapy* became available, which supplies recombinant glucocerebrosidase. This treatment is effective but costs about \$550,000 a year, and takes several hours to infuse each week.

In 2003 came a different approach: *substrate reduction therapy*. This is a drug taken by mouth that decreases the amounts of the substrate, the molecule on which the deficient enzyme acts. A still-experimental third approach is called *pharmacological chaperone therapy*, in which an oral drug binds to a patient's misfolded enzyme, stabilizing it sufficiently to allow some function. Two existing drugs used to treat heart disease act in this way, restoring protein folding in cells from Gaucher's patients.

Gene Therapy

More than a thousand clinical trials of gene therapies have been conducted since 1990. As the new millennium dawned, researchers had expected that the sequencing of the human genome would accelerate the pace of gene therapy development. Instead, new information about the complexity of how genes interact, and a few cases where the experimental treatment harmed the patient, have led to a reevaluation of the idea that we can augment or replace a gene with predictable effects.

Altering genes to treat an inherited disorder theoretically can provide a longer-lasting effect than treating symptoms, but this is much easier said than done. The first gene therapy efforts focused on inherited disorders that researchers knew the most about, even though the conditions are very rare, because these would be the easiest to work with. Gene therapy efforts are now targeting more common illnesses, such as heart disease and cancers. **Tables 20.4** and **20.5** list some general requirements and concerns related to gene therapy.

Types and Targets of Gene Therapy

Gene therapy approaches vary in the way that healing genes are delivered and to which cells they are sent.

Germline gene therapy alters the DNA of a gamete or fertilized ovum. As a result, all cells of the individual have the change. Germline gene therapy is heritable—it passes to

Table 20.4 Gene Therapy Concerns

Scientific	Bioethical
1. Which cells should be treated, and how?	1. Does the participant in a gene therapy trial truly understand the risks?
2. What proportion of the targeted cell population must be corrected to alleviate or halt progression of symptoms?	2. If a gene therapy is effective, how will recipients be selected, assuming it is expensive at first?
3. Is overexpression of the therapeutic gene dangerous?	3. Should rare or more common disorders be the focus of gene therapy research and clinical trials?
4. Is it dangerous if the altered gene enters cells other than the intended ones?	4. What effect should deaths among volunteers have on research efforts?
5. How long will the affected cells function?	5. Should clinical trials be halted if the delivered gene enters the germline?
6. Will the immune system attack the introduced cells?	
7. Is the targeted DNA sequence in more than one gene?	

offspring. It is not being done in humans, although it creates the transgenic organisms discussed in chapter 19.

Somatic gene therapy corrects only the cells that an illness affects. It is nonheritable: A recipient does not pass the genetic correction to offspring. Clearing lungs congested from cystic fibrosis with a nasal spray containing functional CFTR genes is a type of somatic gene therapy.

Gene therapy approaches vary in invasiveness (**figure 20.5**). Cells can be altered outside the body and then infused, called **ex vivo gene therapy**. In **in situ gene therapy**, the functional gene plus the DNA that delivers it (the vector) are injected into a very localized and accessible body part, such as a single melanoma skin cancer. In the most invasive approach, **in vivo** (“in the living body”) **gene therapy**, the gene and vector are introduced directly into the body.

Researchers obtain therapeutic genes using the recombinant DNA and polymerase chain reaction technologies described in chapter 19. In the future, researchers and, someday, clinicians may deliver “artificial” genes synthesized on microchips.

The physical, chemical, and biological methods discussed in section 19.3 are used to send DNA into cells. Physical methods include electroporation, microinjection, and particle

Table 20.5

Requirements for Approval of Clinical Trials for Gene Therapy

1. Knowledge of defect and how it causes symptoms
2. An animal model
3. Success in human cells growing *in vitro*
4. No alternate therapies, or patients for whom existing therapies are not possible or have not worked
5. Safe experiments

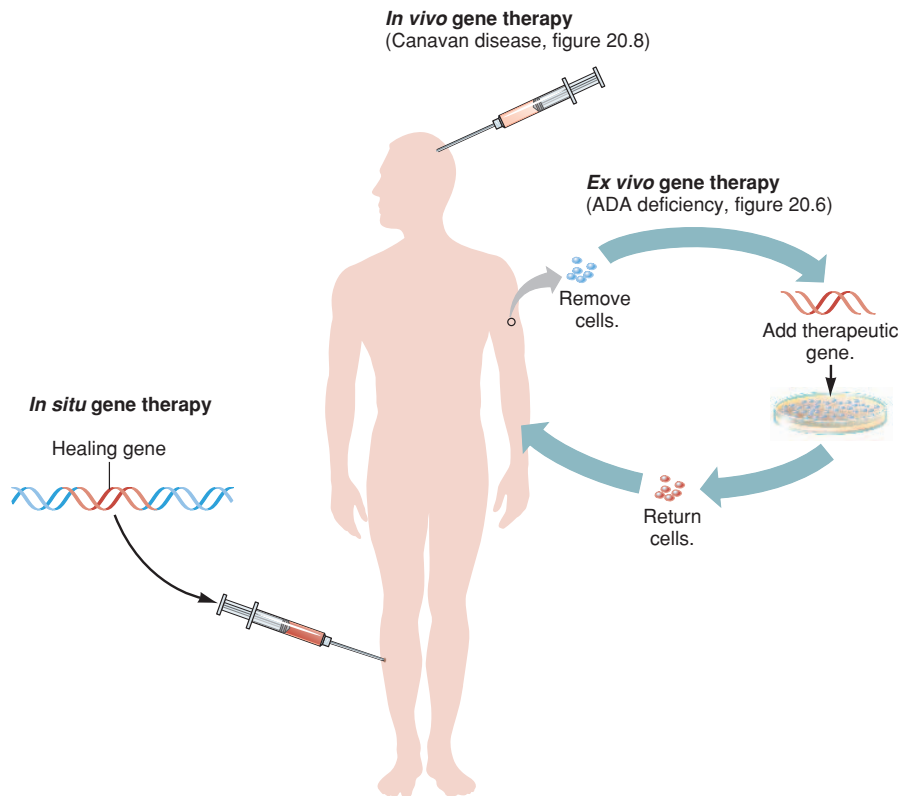


Figure 20.5 Gene therapy invasiveness. Therapeutic genes are delivered to cells removed from the body that are then returned (*ex vivo* gene therapy); delivered directly to an accessible body part such as skin (*in situ* gene therapy); or delivered directly to an interior body part, such as through the skull for Canavan disease or to an artery leading to the liver (*in vivo* gene therapy).

bombardment. Chemical methods include liposomes and other types of lipids that carry DNA across the plasma membrane. The lipid carrier can penetrate the plasma membrane that DNA alone cannot cross, but it may not deliver a sufficient payload, and gene expression is only temporary.

Biological approaches to gene transfer use a vector, such as a viral genome. Researchers remove the viral genes that cause symptoms or alert the immune system and add the corrective gene. Different viral vectors are useful for different types of experiments. A certain virus may transfer its cargo with great efficiency to a specific cell type but carry only a short DNA sequence. Another virus might carry a large piece of DNA but enter many cell types, causing side effects. Or a virus may not infect enough cells to alleviate symptoms. Some retroviruses have limited use because they infect only dividing cells.

Some gene therapies can use viruses that normally infect the targeted cells. For example, adenoviruses that transport CFTR genes to the airway passages of people with cystic fibrosis normally infect lung tissue. A herpes simplex virus delivers the gene encoding enkephalin, a pain-relieving peptide, to nerve endings in skin. Researchers can combine parts of viruses to target a certain cell type. Adeno-associated virus (AAV), for example, infects many cell types, but adding

a promoter from a parvovirus gene restricts it to red blood cell progenitors in bone marrow. Add a human gene that encodes a protein normally found in red blood cells, and the entire vector can treat an inherited disorder of blood, such as sickle cell disease.

Somatic gene therapies are in clinical trials, targeting several different tissues. Gene delivery may be directly to the affected tissue, or into cells that can produce the needed protein and divide. Researchers are increasingly turning to stem and progenitor cells for gene therapies, because these cells can divide as well as travel.

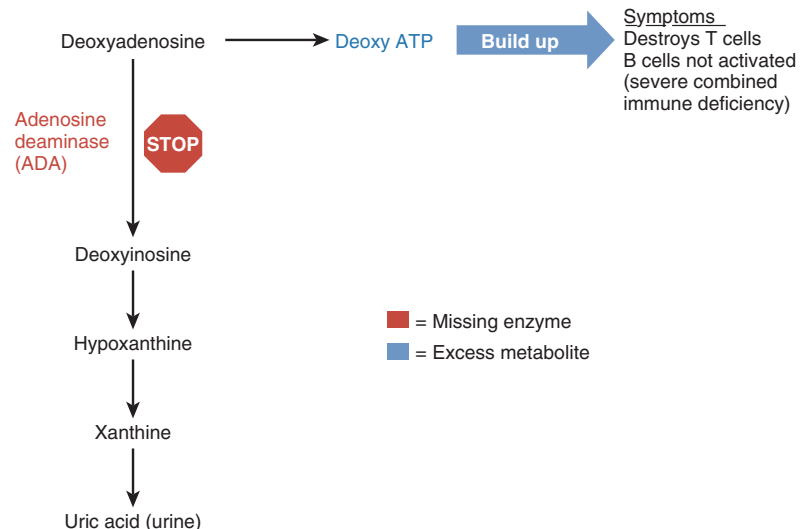
Following are descriptions of a few somatic gene therapies under investigation.

Endothelium Endothelium forms capillaries, which are the tiniest blood vessels. Genetically altered endothelium can secrete a needed protein directly into the bloodstream.

Skin Skin cells grow well. A person can donate a patch of skin the size of a letter on this page; after a genetic manipulation, the sample can grow to the size of a bathmat within 3 weeks, and the skin can be grafted back onto the person. Skin grafts can be genetically modified to secrete therapeutic proteins.



a.



b.

Figure 20.6 Correcting ADA deficiency. (a) Laura Cay Boren spent much of her life in hospitals until she received the enzyme that her body lacks. Here, she pretends to inject her doll as her mother looks on. Today, gene therapy is possible using cord blood stem cells. (b) ADA deficiency causes deoxy ATP to build up, destroying T cells, which therefore cannot stimulate B cells to secrete antibodies. The result is severe combined immune deficiency (SCID).

Muscle Muscle tissue is a good target for gene therapy because it comprises about half of the body's mass, is easily accessible, and is near a blood supply. Several types of cells in muscles are being investigated as treatments for muscular dystrophies, in mice, dogs, and humans. Enough cells must correct the problem to improve muscle strength, and their production of dystrophin must persist.

Liver This largest organ is an important candidate for gene therapy because it has many functions and can regenerate. To treat some inborn errors, as little as 5 percent of the liver's 10 trillion cells would need to be corrected.

Lungs The respiratory tract is easily accessed with an aerosol spray, eliminating the need to remove, treat, and reimplant cells. Several aerosols to treat cystic fibrosis replace the defective gene, but the correction is short-lived and localized.

Nervous Tissue Neurons are difficult targets because they do not divide. Gene therapy can alter other cell types, such as fibroblasts to secrete nerve growth factors or manufacture the enzymes necessary to produce certain neurotransmitters. Then the altered cells are implanted.

Cancer About half of current gene therapy trials target cancer. These approaches enable cancer cells, or their neighbors, to produce proteins that dampen oncogene expression, bolster tumor suppression, strengthen or redirect the immune response, or induce apoptosis.

The Pioneers: Inherited Immune Deficiency

Any new medical treatment begins with courageous volunteers who know that they may risk their health. Gene therapy, however, is unlike conventional drug therapy in that it attempts to alter an individual's genotype in a part of the body that has malfunctioned. Because the potentially therapeutic gene is usually delivered with other DNA, and it may be taken up by cell types other than those affected in the disease, reactions are unpredictable. Following is a look at some of the pioneers of gene therapy.

For the first few years of her life, Laura Cay Boren didn't know what it was like to feel well (**figure 20.6a**). From her birth in July 1982, she fought infection. Colds rapidly became pneumonia, and routine vaccines caused severe abscesses. In February 1983, doctors identified Laura's problem—severe combined immune deficiency (SCID) due to adenosine deaminase (ADA) deficiency (MIM 102700).

Lack of ADA blocks a biochemical pathway that normally breaks down a metabolic toxin into uric acid, which is then excreted. The substance that ADA normally acts upon builds up and destroys T cells. Without helper T cells to stimulate B cells, no antibodies are made. The child becomes very prone to infections and cancer, and usually does not live beyond a year in the outside environment.

The Duke University Medical Center, where Laura celebrated her first and second birthdays, became her second home. In 1983 and 1984, she received bone marrow transplants from her father, which temporarily bolstered her immunity. Red blood cell transfusions also helped, but Laura was still spending more time in the hospital than out. By the end of 1985, she was gravely ill. She had to be fed through a tube,

and repeated infection had severely damaged her lungs. Then Laura was chosen to participate in a trial for a new treatment, and in the spring of 1986, she received her first injection of PEG-ADA. This is the missing enzyme, ADA, from a cow and stabilized with polyethylene glycol (PEG) chains. (PEG is the major ingredient in antifreeze.)

Previous enzyme replacement without PEG didn't work, because what remained of the immune system destroyed the injected, unaltered enzyme. Patients needed frequent doses, which provoked the immune system further, causing severe allergic reactions. Laura's physicians hoped that adding PEG would keep ADA in her blood long enough to work.

Laura began responding to PEG-ADA almost immediately. After 3 months of treatment, toxins no longer showed up in her blood, but her immunity was still suppressed. After 6 months, though, Laura's immune function neared normal for the first time ever—and stayed that way, with weekly doses of PEG-ADA. Her life changed dramatically as she ventured beyond the hospital's germ-free rooms. By summer 1988, she could finally play with other children without fear of infection. She began first grade in fall 1989, but had to repeat the year—she had spent her time socializing! Sadly,

Laura Cay passed away shortly before her 19th birthday, from lung damage.

PEG-ADA revolutionized treatment of this form of SCID, but it replaced the protein, not the gene. The gene therapy approach began on September 14, 1990, at 12:52 P.M. Four-year-old Ashanthi DeSilva sat up in bed at the National Institute of Health in Bethesda, Maryland, and began receiving her own white blood cells intravenously. Earlier, doctors had removed the cells and inserted functioning ADA genes. The gene delivery worked, but did not alter enough cells to restore immunity. It had to be repeated, or PEG-ADA given at intervals. However, Ashanthi is now healthy, and she tells her story at scientific meetings (see *In Their Own Words* below).

A longer-lasting treatment altered progenitor T cells, which account for only one in several billion bone marrow cells, but are more plentiful in umbilical cord blood. If fetuses who had inherited ADA deficiency could be identified, then stem cells could be extracted from their cord blood at birth, given ADA genes, and reinfused.

Crystal and Leonard Gobeia had already lost a 5-month-old baby to ADA deficiency when amniocentesis revealed that their second fetus was affected. They and two other couples

In Their Own Words

The First Gene Therapy Patient

In the late 1980s, the DeSilvas did not think their little girl, Ashanthi ("Ashi"), would survive. She suffered near-continual coughs and colds, and was so fatigued that she could walk only a few steps before becoming winded, her father Raj recalls. "We took her to so many doctors that I stopped counting. One doctor after another would say it was asthma, an allergy, or bronchitis."

Raj's brother, an immunologist, suggested the blood tests that would eventually reveal Ashi's underlying problem—severe combined immune deficiency due to adenosine deaminase (ADA) deficiency. Although unlucky in inheriting a disease, Ashi was lucky in that it was a condition so well understood that it was first in line for gene therapy. Through a series of physician contacts, Ashi became the first recipient.

The medical team at the National Institute (now Institutes) of Health—W. French Anderson, Kenneth Culver, and Michael Blaese—had spent years planning the gene therapy, and were fairly certain that it would work. Within weeks following the therapy, Ashi began to make her own, functional T cells. Although she required further treatments, today she is well and excited about pursuing a career in the music industry.

Over the years, she has championed gene therapy at biomedical conferences. The photo shows her at a meeting when she was 17, where she introduced Dr. Blaese: "Our duty on Earth is to help others. I thank you from the bottom of my heart for all you have enabled me to do."

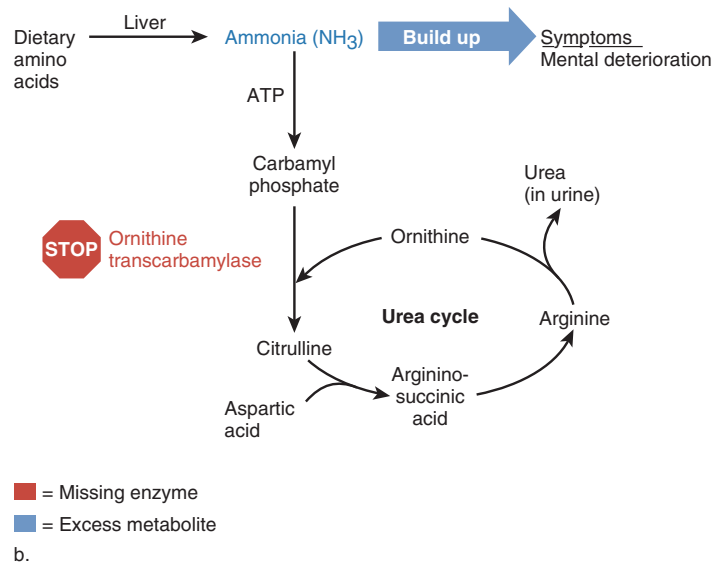
Gene therapy has hit snags in recent years, but overall has had an excellent track record. Said Dr. Blaese a few years ago, "You

have to consider the context. In the years since the first patient, there has been one death and two malignancies. Compare that to the first 100 heart transplants, where only one person lived more than a year. Gene therapy has had a remarkable safety record, yet there are still problems."





a.



b.

Figure 20.7 A brave example. (a) Jesse Gelsinger received gene therapy for an inborn error of metabolism in September 1999. He died four days later from an overwhelming immune response. (b) Lack of an enzyme causes ammonia to accumulate, which is toxic to the brain.

participated in an experiment. Andrew Goba and the other two babies received their own bolstered cord blood cells on the fourth day after birth, with PEG-ADA to prevent symptoms in case the gene therapy did not work right away. T cells carrying normal ADA genes gradually appeared in their blood. By the summer of 1995, the three toddlers each had about 3 in 100 T cells carrying the ADA gene, and they continued to improve.

A few years after the three children with ADA deficiency were treated, another gene therapy trial for SCID began in France. Nine baby boys with a type of X-linked SCID had T cell progenitors removed and given the gene they were missing, which encodes part of a cytokine receptor. The therapy worked, but caused leukemia in three boys when the retrovirus that delivered the therapeutic gene inserted into a proto-oncogene. The boys were successfully treated for the leukemia, but this very unexpected side effect initially stalled many gene therapy trials. It was not the first time such an experiment had a tragic outcome. First came Jesse Gelsinger.

A Major Setback

Eighteen-year-old Jesse Gelsinger died in September 1999, days after receiving gene therapy. An overwhelming immune response to the DNA used to introduce the therapeutic gene killed him.

Jesse had ornithine transcarbamylase deficiency (OTC) (MIM 311250). In this X-linked recessive disorder, one of five enzymes required to break down amino acids liberated from dietary proteins is absent (**figure 20.7**). The nitrogen released from the amino acids combines with hydrogens to form ammonia (NH_3), which rapidly accumulates in the bloodstream and travels to the brain, with devastating effects. The condition

usually causes irreversible coma within 72 hours of birth. Half of affected babies die within a month, and another quarter by age 5. The survivors can control their symptoms by following a special low-protein diet and taking drugs that bind ammonia.

Jesse wasn't diagnosed until he was 2, because he was a mosaic—some of his cells could produce the enzyme, so his symptoms were milder. When he went into a coma in December 1998 after missing a few days of his medications, he and his father considered volunteering for a gene therapy trial they had read about. When he turned 18, Jesse was tested at the University of Pennsylvania, and admitted to the trial. He was jubilant. He knew he might not directly benefit, but he had wanted to try to help babies who die of the condition. A bioethics committee had advised that the experimental treatment not be tried on newborns because the parents would be too distraught to give informed consent. Instead, volunteers were older affected males and carrier females. Said Jesse at the time, "What's the worst that can happen to me? I die, and it's for the babies."

The gene therapy was an adenovirus with a functional human *OTC* gene inserted. This virus had already been used in many gene therapy experiments and did not have the genes to replicate and cause respiratory symptoms. Seventeen patients had already been treated, without side effects, when Jesse entered the hospital to receive a few billion altered viruses in an artery leading into his liver. That night, Jesse developed a high fever. By morning, the whites of his eyes were yellow, indicating that his liver was struggling to dismantle the hemoglobin released from burst red blood cells. A flood of hemoglobin meant a flood of protein, so the ammonia level in his liver skyrocketed, reaching ten times normal levels by mid-afternoon. Jesse became disoriented, then comatose. His lungs and then other vital organs began to fail, and 5 days after beginning

treatment, he was brain dead. His dedicated and devastated medical team stood by as his father turned off life support, and Jesse died.

The autopsy showed that Jesse had had a parvovirus infection, which may have led his immune system to attack the adenovirus. In the liver, the adenovirus had targeted not the hepatocytes as expected, but macrophages that function as sentries for the immune system. In response, interleukins flooded his body, and inflammation raged. Although his parents of children with OTC implored government officials to continue to fund the research, the death of Jesse Gelsinger led to suspension of several gene therapy trials. The death drew particular attention to safety because, unlike most other volunteers, Jesse had not been very ill.

Two Success Stories

Efforts begun in 1995 to treat Canavan disease continued, despite Jesse Gelsinger's fate. Canavan disease is an ideal candidate for gene therapy for several reasons:

1. The gene and protein are well known.
2. There is a window of time when affected children are healthy enough to be treated.
3. Only the brain is affected.
4. Brain scans can monitor response to experimental treatment.
5. No traditional treatment exists.

Canavan disease disrupts the interaction between neurons and neighboring cells called oligodendrocytes, which produce the fatty myelin that coats neurons, enabling them to transmit impulses fast enough for the brain to function (**figure 20.8**). Specifically, the brain neurons normally release N-acetylaspartate (NAA), which is broken down into harmless compounds by an enzyme, aspartoacylase, that the oligodendrocytes produce. In Canavan disease, the enzyme is missing, and the resulting NAA buildup destroys the oligodendrocytes. Without sufficient myelin, neurons cease to function, and developmental delay occurs, as happened to Max Randell, described in the chapter opening essay. Due to a powerful founder effect, Canavan disease is seen almost exclusively in the Ashkenazi Jewish population. *Bioethics: Choices for the Future* on page 409 relates the bitter battle over access to genetic tests for Canavan disease.

The first attempts at gene therapy for Canavan disease introduced the gene in a liposome, through holes bored into the skull. The first recipient, an 18-month-old, regained some skills for awhile. Previously, she could barely open her eyes and did not interact with anyone. But 3 months after the therapy, she looked around, moved, and vocalized. A brain scan showed neuron myelination in regions where it had

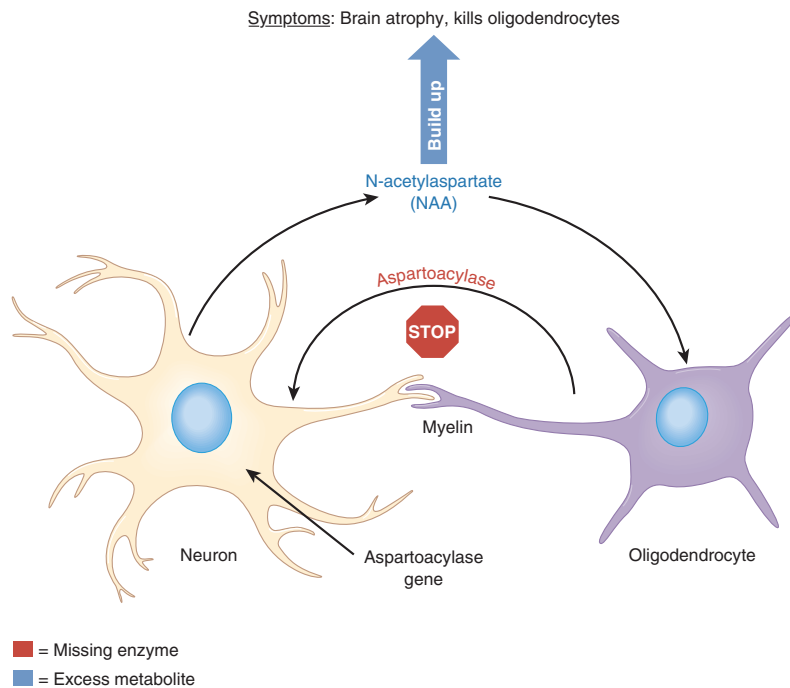


Figure 20.8 Canavan disease. Stripping of the lipid layer on brain neurons occurs because oligodendrocytes lack an enzyme to break down NAA, which neurons produce. Gene therapy enables brain neurons to secrete the enzyme, restoring the fatty covering that makes nerve transmission possible.

vanished. She was not treated again until June 2001, when a viral vector replaced the liposomes. In the interim, while regulatory agencies argued about the safety of this therapy for a disease that had no other treatment, she lost some of the gains from the first round, such as being able to hold her head up. But the gene therapy does appear to be working in other children.

The second gene therapy success story is for an autosomal recessive form of blindness called Leber's congenital amaurosis II (MIM 204100). Although the disorder is the most severe inherited form of blindness, present from birth, several months may pass before parents realize that their infant's lack of eye contact signals blindness. An affected child can sense light and may suffer night blindness, but usually isn't completely sightless until the third decade. The rods and cones, which are photoreceptors in the retina at the back of the eye, degenerate. Rods provide black and white vision, and cones provide color vision.

A mutation in a gene called *RPE64* causes Leber's congenital amaurosis II. "RPE" stands for retinal pigment epithelium, referring to the cells that provide the pigment rhodopsin to the rods and cones. Gene therapy for the disorder began with a natural animal model for this form of blindness—a breed of dog called a Briard that has the same mutation and phenotype as humans (**figure 20.9a**). In 1998, researchers injected the wild type alleles for the gene

Bioethics: Choices for the Future

Canavan Disease: Patients Versus Patents

When Debbie Greenberg gave birth to Jonathan in 1981, she and her husband Dan had no idea that they would one day lead an effort to challenge how a researcher and a hospital patented a gene. Jonathan lived 11 years with Canavan disease; his brain never developed past infancy. The couple had an affected daughter, Amy, a few years after Jonathan was born, and three healthy children.

Shortly after Jonathan's diagnosis, the Greenbergs started the Canavan Foundation, which established a tissue bank that stored blood, urine, and autopsy tissue from affected children. In 1987, the Greenbergs met Dr. Reuben Matalon at a Tay-Sachs disease screening event in Chicago, and convinced him to search for the Canavan gene. The Greenbergs helped to collect tissue from families from all over the world, which was critical to Dr. Matalon's success in identifying the gene and the causative mutation in 1993, when he was working at Miami Children's Hospital.

Detecting the mutation could be used to confirm diagnoses, detect carriers, and test for the condition prenatally. By 1996, the Canavan Foundation was offering free testing. But unknown to the members of the organization who had donated their children's tissues for the gene search, Dr. Matalon and Miami Children's Hospital had filed for a patent on their discovery.

The U.S. Patent and Trademark office granted invention number 5,679,635—the Canavan gene—in 1997. A year later, the American College of Obstetricians and Gynecologists advised their physician members to offer carrier testing to Ashkenazi Jewish patients, because one in forty such women is a carrier. Identifying couples in which both people are carriers would give them the option of avoiding giving birth

to affected children. That same year, Miami Children's Hospital began to exercise its patent rights by requiring that doctors and diagnostic laboratories charge for a Canavan test. Suddenly, families whose donations—both monetary and biological—had made the discovery of the gene possible had to pay for carrier and prenatal tests. They were outraged.

On November 30, 2000, a group of parents and three nonprofit organizations filed suit in Chicago against Dr. Reuben Matalon and Miami Children's Hospital. The suit does not challenge the patent, but how it was obtained—in secret, they claim. They wish to recover earnings from the gene test to be turned over to the families who had to pay to offset licensing fees. In 2003, U.S. district judge Federico Morena upheld the rights of the parents to sue, calling the case “a tale of successful research collaboration gone sour.”

Questions for Discussion

1. It is common today for families with genetic diseases to start organizations that help researchers better understand the disease. Go to the Genetic Alliance or National Organization of Rare Diseases or other websites and compare and contrast how different organizations have learned to protect themselves from having to pay for tests that they helped to develop.
2. Which organizations or individuals should provide oversight to the use of patients' cells and tissues in developing medical tests?
3. What is a justification for a pharmaceutical company charging families for tests that their biological materials helped to develop?

aboard a viral vector into the retinas of several dogs. They regained their eyesight, and one animal, Lancelot, even appeared before Congress to lobby for increased funding for gene therapy.

After the success of the gene therapy on dogs, the next step was to try the approach on people. The first experiments were done on four young adults. The trial was planned to test only safety, with a single, low dose. The treatment was not only apparently safe, but improved the participants' vision! They could detect more light, could read part of an eye chart, and navigate an obstacle course. Next, the gene therapy was tried on an 8-year-old from Lake Luzerne, New York (figure 20.9b). Before his therapy, young Corey had night blindness, extreme nearsightedness, and a visual field so narrow that his view of the world was like looking through a tiny tunnel. At school he used a computer screen to display the chalkboard, plus a magnifying glass. Shortly after his first treatment, on a sunny day, Corey's parents took him to the zoo. They noticed that his eyes seemed bluer, and he complained of the light intensity. Corey's pupils were finally reacting to light! Faces soon began to emerge from his hazy vision, and he continues to

improve. Ironically, the successful gene therapy trials for Leber's congenital amaurosis are being done at the Children's Hospital of Philadelphia—the same city where Jesse Gelsinger's gene therapy failed.

Expectations and Limitations

When the age of gene therapy dawned in the 1990s, expectations were high—and for good reason. Work in the 1980s had clearly shown abundant, pure, human biochemicals, useful as drugs, could come from genetically modified cells and transgenic organisms. It was a matter of time, many people thought, before genetic altering of our own somatic tissue would treat a variety of ills.

In reality, gene therapy progress has been slow. Boys with Duchenne muscular dystrophy who receive immature muscle cells with healthy dystrophin genes do not walk again, although they might be able to wiggle a toe. People with cystic fibrosis who inhale viruses bearing the CFTR gene do not permanently breathe easier, but might feel relief for a few weeks. Others have died since Jesse Gelsinger.



a.



b.

Figure 20.9 A gene therapy success story in-the-making. Gene therapy for Leber's congenital amaurosis II was tried first on Briard dogs (a), then on four young adults, and then on 8-year-old Corey Haas (b). So far it is working!

The sequencing of the human genome did not provide a list of new gene defects to correct—many of the disease-causing genes were already known—but instead revealed a complexity to genome structure and function that will impact gene therapy. Consider the fact that the same exon sequence can be part of different genes. Targeting an exon because it is part of one gene may affect others, healing one set of symptoms while causing others. As genome researchers continue to identify gene functions, this risk should lessen.

Discovery of microRNAs, discussed in chapter 11, may also complicate gene therapy. Correcting a genetic mistake in the nucleus may not counter a disease phenotype because of what may happen in the cytoplasm—the mRNA transcribed from the delivered gene may be silenced before the needed protein is synthesized. Yet another area of uncertainty is the issue of somatic versus germline gene therapy. A corrected gene targeted to a particular tissue may find its way, in the circulation, to the reproductive tract, enter a gamete, and thereby affect the next generation.

Despite these drawbacks—real and theoretical—at a molecular and cellular level, gene therapy *is* working. The patients with muscular dystrophy, cystic fibrosis, and SCID have

cells that have accepted and expressed therapeutic genes. The challenge now is to find just the right vectors to deliver sustained, targeted, and safe genetic corrections.

Key Concepts

1. Protein-based therapies replace gene products and treat the phenotype.
2. Gene therapies replace malfunctioning or absent genes.
3. Germline gene therapy targets gametes or fertilized ova and is heritable. Somatic gene therapy targets various types of somatic tissue as well as cancer cells and is not heritable.
4. Gene therapy may be *in situ*, *ex vivo*, or *in vivo*.
5. Vectors for delivering genes include liposomes and viruses.
6. Gene therapies have targeted ADA deficiency, OTC deficiency, Canavan disease, and a form of hereditary blindness.

Summary

20.1 Geneticists Find Zebras, and some Horses

1. Geneticists are skilled at seeing the single-gene disorders that underlie specific sets of signs and symptoms.

20.2 Genetic Counseling

2. **Genetic counselors** provide information on inheritance patterns, disease risks and symptoms, and available tests and treatments.
3. Prenatal counseling and counseling a family coping with a particular disease pose different challenges.
4. Genetic counselors interpret direct-to-consumer genetic tests and assist other health care professionals in incorporating genetic information into their practices.

20.3 Genetic Testing

5. Newborns are routinely screened for several inborn errors of metabolism, some of which are treatable. The numbers of tests vary by state.
6. The Clinical Laboratory Improvement Amendments regulate some genetic tests. Direct-to-consumer tests presented as information and not diagnoses may not be regulated.
7. Nutrigenetics tests may provide inaccurate information.
8. **Pharmacogenetics** and **pharmacogenomics** provide information on how individuals metabolize certain drugs.
9. Enzyme replacement therapy, substrate reduction therapy, and pharmacological chaperone therapy are approaches to treating the biochemical imbalances of certain inborn errors.

20.4 Treating Genetic Disease

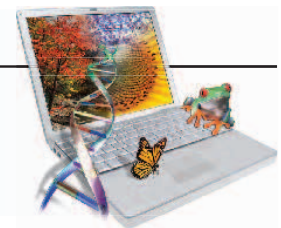
10. **Germline gene therapy** affects gametes or fertilized ova, affects all cells of an individual, and is transmitted to future generations. It is not performed in humans. **Somatic gene therapy** affects somatic tissue and is not passed to offspring.
11. **Ex vivo gene therapy** is applied to cells outside the body that are then reimplanted or reinfused into the patient. **In situ gene therapy** occurs directly on accessible body parts. **In vivo gene therapy** is applied in the body.
12. Gene therapy delivers new genes and encourages production of a needed substance at appropriate times and in therapeutic (not toxic) amounts.
13. Several types of vectors are used to deliver therapeutic genes. Some gene therapies target stem or progenitor cells, because they can divide and move.
14. Development of gene therapy has been slower than anticipated because of the unexpected complexities of gene interactions and the challenge of adequately targeting and sustaining therapeutic effects.

Review Questions

1. What is unique about the services that a genetic counselor provides compared to those of a nurse or physician?
2. What are the advantages and disadvantages of “virtual” genetic counseling (using an interactive computer program rather than a human)?
3. Why is newborn screening economically feasible?
4. Using information from this or other chapters, or the Internet, cite genetic tests given to a newborn, young adult, and middle-aged person (three different tests).
5. Select a disease or health problem, such as depression or cancer, and explain how a pharmacogenetic or pharmacogenomic approach to drug treatment can be more effective and tolerable than the older approach that treats patients with the same disorder as if they will respond in the same way to a particular drug.
6. What are the three stages of the evolution of treatments for single-gene disorders?
7. Why is the removal of blood in people with hereditary hemochromatosis not gene therapy?
8. Explain the differences among *ex vivo*, *in situ*, and *in vivo* gene therapies. Give an example of each.
9. Would somatic gene therapy or germline gene therapy have the potential to affect evolution? Explain your answer.
10. What factors would a researcher consider in selecting a viral vector for gene therapy?
11. What are some of the complications that have slowed the development of gene therapy?
12. Compare the risks and potential benefits of gene therapy.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Applied Questions

1. Discuss the challenges that a genetic counselor faces in explaining to parents-to-be a prenatal diagnosis of trisomy 21 in a fetus with no family history, compared to a prenatal diagnosis of translocation Down syndrome in a family with a reproductive history of pregnancy loss and birth defects. (Chapter 13 discusses Down syndrome.)
2. A company tests for variants of the gene *ACTN3* (alpha-actinin 3), which encodes a protein that binds actin, a cytoskeletal protein. One genotype is more common among elite sprint athletes, and another among endurance athletes. Some parents are testing their young children for these gene variants and using the results to decide whether the child should pursue a sport that entails sprinting or endurance. Opponents of the test point out that simply having kids run a race would provide more meaningful information than would data on this single gene. Would you have your child tested? Cite a reason for your answer.
3. In the past, when invasive prenatal tests were less safe, health care providers typically told only women over age 35 or with a family history of Down syndrome about these tests, because given the technology of the time, they were at higher risk than other women. Since the safety has improved, more women are being offered prenatal diagnosis of Down syndrome. In some nations, doctors are now required to offer it to all pregnant women. Some women have successfully sued for “wrongful births” of children with Down syndrome because their doctors failed to tell them of or refer them for prenatal testing. *Bioethics: Choices for the Future* on page 245 provides further information.

Discuss the idea of a “wrongful birth” from the point of view of the parent suing the doctor, the child with Down syndrome, the doctor, and a family support group for individuals with Down syndrome.

4. The Newborn Screening Translational Research Network Coordinating Center is a new program to coordinate newborn screening among states. It plans to analyze stored bloodspots for information on treatments and outcomes. Do you think that the parents of the newborns whose blood is being analyzed in new ways should be consulted, or is their consent presumed because newborn screening is mandated by state law?
5. Choose one of the cases described in the chapter, and write out how you would counsel the family after the initial diagnosis.
6. Why would the American College of Medical Genetics ask that the government mandate testing for inborn errors of metabolism that do not have treatments?
7. Create a gene therapy by combining items from the three lists below. Describe the condition to be treated, and how a gene therapy might correct the symptoms.
8. Genes can be transferred into the cells that form hair follicles. Would gene therapy to treat baldness most likely be *ex vivo*, *in situ*, or *in vivo*? Cite a reason for your answer.

Cell Type	Vector	Disease Target
fibroblast	AV	Duchenne muscular dystrophy
skin cell	AAV	Alzheimer disease
neuroglial cell	retrovirus	sickle cell disease
red blood cell	liposome	cystic fibrosis
progenitor cell		
myoblast		

Web Activities

9. Consult any of these websites to determine the number of newborn screening tests in your state:
 - <http://www.huntershpe.org/UNBS/index.html>
 - marchofdimmes.com/peristats
 - genes-r-us.uthscsa.edu
 Select one of the diseases, then consult the websites for Genetic Alliance or the National Organization for Rare Diseases, and describe the symptoms and treatments.
10. Go to <http://www.pharmgkb.org/> and click on VIP. Select a gene and discuss its importance in health care.
11. Go to www.genedx.com. Describe a genetic test that this company offers.
12. Go to www.medomics.com and click on Dx Genome. Read this organization's goals. State two ways that their approach differs from direct-to-consumer websites such as www.23andme.com

Case Studies and Research Results

13. How would you, as a genetic counselor, handle the following situations (all real)? What would you tell the patients, and what tests would you suggest? (See other chapters for specific information.)
 - a. A couple in their early forties is expecting their first child. Amniocentesis indicates that the fetus is XXX, which might never have been noticed without the test. When they learn of the abnormality, the couple asks to terminate the pregnancy.
 - b. Two people of normal height have a child with achondroplastic dwarfism, an autosomal dominant trait. They are concerned that subsequent children will also have the condition.
 - c. A newborn has a medical condition not associated with any known gene mutation or chromosomal aberration. The parents want to sue the genetics department of the medical center because the amniocentesis did not indicate a problem.
14. Three-year-old Tawny Fitzgerald has been to the emergency department repeatedly for broken bones. At the last visit, a nurse questioned Tawny's parents, Donald and Rebecca, about possible child abuse. No charges were filed—the child just appeared to be clumsy. Then Tawny's brother Winston was born. When he was 6 months old, Donald found him screaming in pain one morning. A trip to the hospital revealed a broken arm. This time, a social worker was sent to the Fitzgerald home. Donald and Rebecca were interviewed in great depth and advised to find a lawyer. A relative in medical school suggested that they have the children examined for osteogenesis imperfecta, also known as “brittle bone disease.” Consult MIM and list the facts about a form of this condition that could affect both sexes, with carrier parents. If you were the genetic counselor hired to help this couple, what would you ask them, and tell them, to help them deal with the legal and social services authorities who might need a biology lesson?
15. Jill and Scott S. had thought 6-month-old Hannah was developing just fine until Scott's sister, a pediatrician, noticed that the baby's abdomen was swollen and hard. Knowing that the underlying enlarged liver and spleen could indicate an inborn error of metabolism, Scott's sister suggested the child undergo several tests. She had inherited sphingomyelin lipidosis, also known as Niemann-Pick disease type A (MIM 257200). Both parents were carriers, but Jill had tested

negative when she took a Jewish genetic disease panel during her pregnancy because her mutation was very rare and not part of the test panel. Hannah was the first child with Niemann-Pick disease to be successfully treated with a transplant of umbilical cord blood cells from a donor. She caught up developmentally and became more alert. Monocytes, a type of white blood cell, from the cord blood traveled to her brain and manufactured the deficient enzyme. Dietary therapy does not work for this condition because the enzyme cannot cross from the blood to the brain. Monocytes, however, can enter the brain.

- a. Did Hannah's treatment alter her phenotype, genotype, or both?
 - b. Why did the transplant have to come from donated cord blood, and not from Hannah's own, which had been stored?
 - c. If you were the genetic counselor, what advice would you give this couple if they conceive again?
16. The Food and Drug Administration has approved tests for variants of two genes that encode cytochrome 450 enzymes that affect the rate at which a person metabolizes selective serotonin reuptake inhibitors, a drug class that includes

widely used antidepressants. However, a study by the Centers for Disease Control found "no evidence . . . showing that the results of CYP450 testing influenced SSRI choice or dose and improved patient outcomes." More than a dozen direct-to-consumer websites offer the tests, several of which claim that the test results can be used to choose the correct antidepressant and dose.

- a. Why might the genotypes be associated with a drug metabolism phenotype, yet not improve outcome?
 - b. Do you think that selling these tests directly to consumers is misleading or unethical?
 - c. Suggest a way to ensure that genetic tests provide useful information.
17. In the mid-1990s, several children with tyrosinemia type I (MIM 276700) were treated with liver transplants. In this inborn error of metabolism, a missing enzyme causes kidney and liver failure. For several years, the transplanted children did very well, but gradually their kidneys began to fail again. Explain why the gene therapy did not completely cure the condition.



Under what circumstances should sperm from a deceased soldier be used to conceive a child?

CHAPTER

21

Reproductive Technologies

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Preimplantation Genetic Diagnosis

21.4 Extra Embryos

Postmortem Sperm Retrieval

Bruce and Gaby V., in their early thirties, had delayed becoming parents, confident that their good health would make pregnancy possible later. But then Bruce suddenly died of an allergic reaction to a medication. Gaby knew how much Bruce had wanted to be a father, so she asked the medical examiner to collect Bruce's sperm. The sample was sent to the California Cryobank, where it lay deeply frozen for more than a year.

In the summer of 1978, Cappy Rothman, medical director of the sperm bank, defrosted the sperm and used it to fertilize one of Gaby's oocytes. On March 17, Bruce and Gaby's daughter was born. It was the first case of postmortem sperm retrieval in which the father did not actively participate in the decision. In another case, the man was dying from cancer and had time to state his wishes. Since 1990, U.S. servicemen who feared infertility from exposure to chemical or biological weapons have taken advantage of sperm bank discounts to the military.

Postmortem sperm retrieval raises legal and ethical issues. A woman conceived twins with her husband's consent 16 months after he died of leukemia at age 30. The Social Security Administration refused to provide survivor benefits to their daughters, claiming that the husband was not a father, but a sperm donor. The Massachusetts Superior Court reversed this decision. Like other assisted reproductive technologies, postmortem sperm retrieval is not regulated at the federal level in the United States. Bioethicists have identified situations to avoid:

- Someone other than a spouse wishing to use the sperm
- A too-hasty decision based on grief
- Use of the sperm for monetary gain

21.1 Savior Siblings And More

A couple in search of an oocyte donor advertises in a college newspaper seeking an attractive young woman from an athletic family. A cancer patient stores her oocytes before undergoing treatment. Two years later, she has several of them fertilized in a laboratory dish with her partner's sperm, and has a cleavage embryo implanted in her uterus. She becomes a mother. A man paralyzed from the waist down has sperm removed and injected into his partner's oocyte. He, too, becomes a parent when he thought he never would.

Lisa and Jack Nash sought to have a child for a different reason. Their daughter Molly, born on July 4, 1994, had Fanconi anemia (MIM 227650). This autosomal recessive condition would destroy her bone marrow and her immunity. An umbilical cord stem cell transplant from a sibling could likely cure her, but Molly had no siblings. Nor did her parents wish to have another child with a one in four chance of inheriting the disorder, as Mendel's first law dictates. Technology offered another solution.

In late 1999, researchers at the Reproductive Genetics Institute at Illinois Medical Center mixed Jack's sperm with Lisa's oocytes in a laboratory dish. After allowing fifteen of the fertilized ova to develop to the eight-cell stage, researchers separated and applied DNA probes to one cell from each embryo. A cell that had wild type Fanconi anemia alleles and that matched Molly's human leukocyte antigen (HLA) type was identified and its seven-celled remainder implanted into Lisa's uterus. Adam was born in late summer. A month later, physicians infused his umbilical cord stem cells into Molly, saving her life (**figure 21.1**).

Increased knowledge of how the genomes of two individuals come together and interact has spawned several novel ways to have children. **Assisted reproductive technologies** (ARTs) replace the source of a male or female gamete, aid fertilization, or provide a uterus. These procedures were developed to treat infertility, but are increasingly encompassing genetic screening. In the United States the government does



Figure 21.1 Savior siblings. Adam Nash was conceived and selected to save his sister Molly's life. He is also a much-loved sibling and son. Several other families have since conceived one child to help another.

not regulate ARTs, but the American Society for Reproductive Medicine provides voluntary guidelines. The United Kingdom has pioneered ARTs and its Human Fertilisation and Embryology Authority has served as a model for government regulation. A great advantage of the British regulation of reproductive health services and technologies is that databases include success rates of the different procedures.

Key Concepts

1. Assisted reproductive technologies provide innovative ways to conceive offspring.
2. ARTs are used to avoid conception of a child with a particular genetic condition, or to overcome infertility.

21.2 Infertility and Subfertility

Infertility is the inability to conceive a child after a year of frequent intercourse without the use of contraceptives. Some specialists use the term *subfertility* to distinguish those individuals and couples who can conceive unaided, but for whom this may take longer than usual. On a more personal level, infertility is a seemingly endless monthly cycle of raised hopes and crushing despair. In addition to declining fertility, as a woman ages, the incidence of pregnancy-related problems rises, including chromosomal anomalies, fetal deaths, premature births, and low-birthweight babies. For most conditions, the man's advanced age does not raise the risk of pregnancy complications, although sperm motility declines with age.

Physicians who specialize in infertility treatment can identify a physical cause in 90 percent of cases. Of these, 30 percent of the time the problem is primarily in the male, and 60 percent of the time it is primarily in the female. When a physical problem is not obvious, the cause is usually a mutation or chromosomal aberration that impairs fertility in the male. The statistics are somewhat unclear, because in 20 percent of the 90 percent, both partners have a medical condition that could contribute to infertility or subfertility. A common combination is a woman with an irregular menstrual cycle and a man with a low sperm count. One in six couples has difficulty in conceiving or giving birth to children.

Male Infertility

Infertility in the male is easier to detect but sometimes harder to treat than female infertility. One in twenty-five men is infertile. Some men have difficulty fathering a child because they produce fewer than the average 20 to 200 million sperm cells per milliliter of ejaculate, a condition called oligospermia. It has several causes. If a low sperm count is due to a hormonal imbalance, administering the appropriate hormones may boost sperm output. Sometimes a man's immune system produces IgA antibodies that cover the sperm and prevent them from binding to

oocytes. Male infertility can also be due to a varicose vein in the scrotum. This enlarged vein produces too much heat near developing sperm, and they cannot mature. Surgery can remove a scrotal varicose vein.

Most cases of male infertility are genetic. About a third of infertile men have small deletions of the Y chromosome that remove the only copies of key genes whose products control spermatogenesis. Other genetic causes of male infertility include mutations in genes that encode androgen receptors or protein fertility hormones, or that regulate sperm development or motility. **Reading 21.1** describes the recently identified first type of autosomal recessive male infertility that is not part of a syndrome.

Sperm with extra chromosomes are ten times more likely to occur in men who had vasectomies reversed. Exactly how this happens isn't known, but it may be related to blocking the male reproductive system to keep developing sperm out of semen. If sperm with an abnormal number of chromosomes fertilize oocytes, the imbalance may end development so early that repeated pregnancy losses appear to be infertility.

For many men with low sperm counts, fatherhood is just a matter of time: They are subfertile, not infertile. If an ejaculate contains at least 60 million sperm cells, fertilization is likely eventually. To speed conception, a man with a low sperm count can donate several semen samples over a period of weeks at a fertility clinic. The samples are kept in cold storage, then pooled. Some of the seminal fluid is withdrawn to leave a sperm cell concentrate, which is then placed in the woman's body. It isn't very romantic, but it is highly effective at achieving pregnancy. Men who actually want a very low sperm count—those who have just had a vasectomy for birth control—can use an at-home test kit to monitor their sperm counts. Fewer than 250,000 sperm cells per milliliter of seminal fluid makes pregnancy highly unlikely.

Sperm quality is more important than quantity. Sperm cells that are unable to move—a common problem—or are shaped abnormally, cannot reach an oocyte. However, the genetic package of an immobile or abnormally shaped sperm cell can be injected into an oocyte, although this does not always lead to fertilization. If the cause of male infertility is hormonal, replacing the absent hormones can sometimes make sperm move. Even sperm that look and move normally may be unable to fertilize an oocyte.

Faulty apoptosis (programmed cell death) can also cause male infertility. Apoptosis normally kills abnormally shaped sperm. Men with high percentages of abnormally shaped sperm often have cell surface molecules that indicate impaired apoptosis.

Female Infertility

Abnormalities in any part of the female reproductive system can cause infertility. (**figure 21.2**). Many women with subfertility or infertility have irregular menstrual cycles, making it difficult to pinpoint when conception is most likely. In an average menstrual cycle of 28 days, ovulation usually occurs around the 14th day after menstruation begins. This is when a woman is most likely to conceive.

For a woman with regular menstrual cycles who is under 30 years old and not using birth control, pregnancy typically happens within 3 or 4 months. A woman with irregular menstrual periods can tell when she is most fertile by using an ovulation predictor test, which detects a peak in the level of luteinizing hormone that precedes ovulation by a few hours. Another way to detect the onset of ovulation is to record body temperature each morning using a digital thermometer with subdivisions of hundredths of a degree Fahrenheit, which can indicate the 0.4 to 0.6 rise in temperature when ovulation starts. Sperm can survive in a woman's body for up to 5 days, but the oocyte is only viable for 24 to 48 hours after ovulation.

The hormonal imbalance that usually underlies irregular ovulation has various causes. These include a tumor in the ovary or in the pituitary gland in the brain that controls the reproductive system, an underactive thyroid gland, or use of steroid-based drugs such as cortisone. Sometimes a woman produces too much prolactin, the hormone that promotes milk production and suppresses ovulation in new mothers. If prolactin is abundant in a nonpregnant woman, she will not ovulate.

Fertility drugs can stimulate ovulation, but they can also cause women to "superovulate," producing more than one oocyte each month. A commonly used drug, clomiphene, raises the chance of having twins from 1 to 2 percent to 4 to 6 percent. If a woman's ovaries are completely inactive or absent (due to a birth defect or surgery), she can become pregnant only if she uses a donor oocyte. Some cases of female infertility are due to "reduced ovarian reserve"—too few oocytes. This is typically discovered when the ovaries do not respond to fertility drugs. Signs of reduced ovarian reserve are an ovary with too few follicles (observed on an ultrasound scan) or elevated levels of follicle-stimulating hormone on the third day of the menstrual cycle.

The uterine tubes are also a common site of female infertility because fertilization usually occurs in open tubes. Blockage can prevent sperm from reaching the oocyte, or entrap a fertilized ovum, keeping it from descending into the uterus. If an embryo begins developing in a blocked tube and is not removed and continues to enlarge, the tube can burst and the woman can die. Such a "tubal pregnancy" is called an ectopic pregnancy.

Uterine tubes can also be blocked due to a birth defect or, more likely, from an infection such as pelvic inflammatory disease. A woman may not know she has blocked uterine tubes until she has difficulty conceiving and medical tests uncover the problem. Surgery can sometimes open blocked uterine tubes.

Excess tissue growing in the uterine lining may make it inhospitable to an embryo. This tissue can include benign tumors called fibroids or areas of thickened lining from a condition called endometriosis. The tissue can grow outside of the uterus too, in the abdominal cavity. In response to the hormonal cues to menstruate, the excess lining bleeds, causing cramps. Endometriosis can hamper conception, but curiously, if a woman with endometriosis conceives, the cramps and bleeding usually disappear after the birth.

Secretions in the vagina and cervix may be hostile to sperm. Cervical mucus that is thick or sticky due to infection can entrap sperm, keeping them from moving far enough to

Reading 21.1

The Case of the Round-Headed Sperm—and a Review of This Book

In fewer than a tenth of a percent of men who are infertile, sperm cells lack the tip, called the acrosome, where the enzymes that break through the layers surrounding an oocyte collect. This condition is called “globozoospermia” (**figure 1**). An Ashkenazi Jewish family led researchers to a gene that, when mutant, causes an autosomal recessive form of male infertility due to round-headed sperm.

The family went to a center for reproductive medicine in Brussels, the Netherlands. Of the six sons, three were infertile (**figure 2**). Three daughters were fertile. The affected sons’ sperm were obviously misshapen, and the mode of inheritance obviously recessive, since the parents were fertile.

Researchers suspected that consanguinity was involved, because a shared ancestor increases risk of inheriting a very rare autosomal recessive condition if the mutation is in the family. But the family denied knowing a relative who had married a relative. Reasoning that perhaps DNA could reveal what the family did not know, researchers scanned the genomes of all six sons for regions of homozygosity. Recall from chapter 7 that these areas in a genome

reflect inheritance from shared ancestors, indicating consanguinity. A region of homozygosity in this case was defined as 25 consecutive SNPs that were homozygous. The genomes of all six sons were riddled with these regions, suggesting that at some point, cousin married cousin or an aunt/uncle wed a nephew/niece. One region of homozygosity was seen in all three infertile brothers, but was heterozygous in two of the three fertile brothers. The remaining brother was homozygous wild type for the region.

Next, the researchers delved into the part of the long arm of chromosome 3 where the telltale region of homozygosity lay. It houses fifty genes, only one of which is expressed specifically in the testes. This gene is called “spermatogenesis-associated 16”, or *SPATA16* (MIM 102530). It has eleven exons, and the mutation in the Ashkenazi family is a single base change, from G to A, at the 848th position in the gene, near the end of exon 4. The mutation likely upsets the splicing out of introns.

The wild type protein product of the *SPATA16* gene is transported from the Golgi apparatus into vesicles that take it to the acrosome as it telescopes out of the front end of a sperm cell. By attaching the gene for the jellyfish’s green fluorescent protein (see figure 19.1) to the wild type *SPATA16* gene in cells growing in culture, researchers visualized the protein being transported to the forming acrosome in immature sperm.



Figure 1 A misshapen sperm cannot fertilize an oocyte.

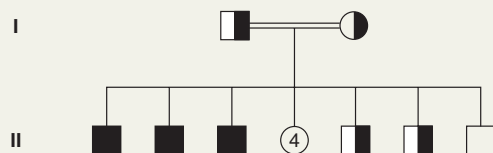


Figure 2 In a family with autosomal recessive globozoospermia, three of six sons are infertile.

encounter an oocyte. Vaginal secretions may be so acidic or alkaline that they weaken or kill sperm. Douching daily with an acidic solution such as acetic acid (vinegar) or an alkaline solution, such as bicarbonate, can alter the pH of the vagina so that in some cases it is more receptive to sperm cells. Too little mucus can prevent conception too; this is treated with low daily doses of oral estrogen. Sometimes mucus in a woman’s body harbors antibodies that attack sperm. Infertility may also result if the oocyte fails to release sperm-attracting biochemicals.

One reason the incidence of female infertility increases with age is that older women are more likely to produce oocytes with an abnormal chromosome number, which often causes spontaneous abortion because defects are too severe for development to proceed for long. The cause is usually misaligned spindle fibers when meiosis resumes, causing aneuploidy (extra

or missing chromosomes). Perhaps the longer exposure of older oocytes to harmful chemicals, viruses, and radiation contributes to the risk of meiotic errors. Losing very early embryos may appear to be infertility because the bleeding accompanying the aborted embryo resembles a heavy menstrual flow.

Infertility Tests

A number of medical tests can identify causes of infertility. The man is checked first, because it is easier, less costly, and less painful to obtain sperm than oocytes.

Sperm are checked for number (sperm count), motility, and morphology (shape). An ejaculate containing up to 40 percent unusual forms is still considered normal, but many more than this can impair fertility. A urologist performs sperm tests.

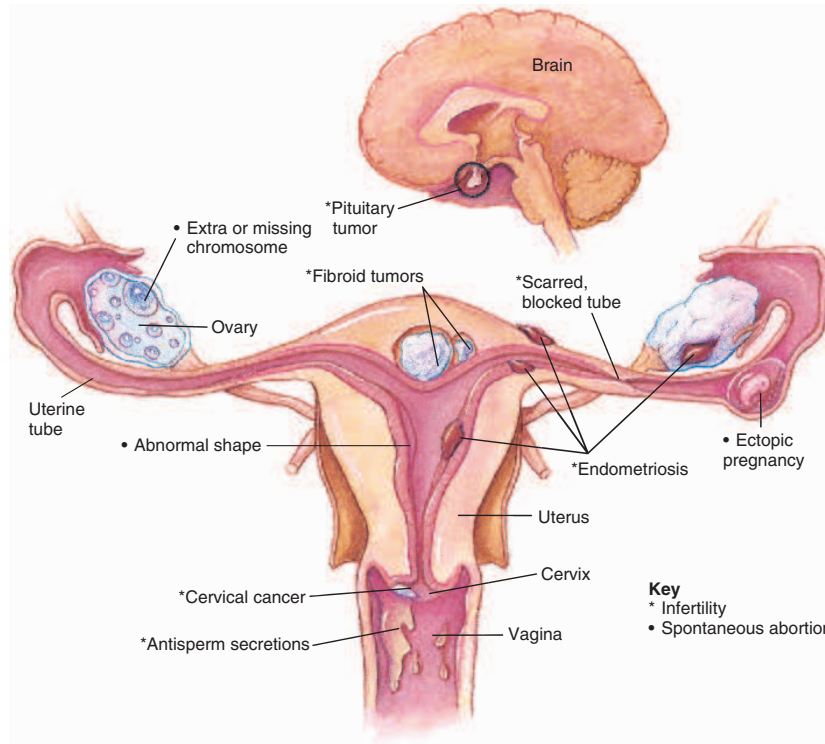


Figure 21.2 Sites of reproductive problems in the female.

A genetic counselor can evaluate Y chromosome deletions associated with lack of sperm. If a male cause of infertility is not apparent, a gynecologist checks the woman to see that reproductive organs are present and functioning.

Some cases of subfertility or infertility have no clear explanation. Psychological factors may be at play, or it may be that inability to conceive results from consistently poor timing. Sometimes a subfertile couple adopts a child, only to conceive one of their own shortly thereafter; many times, infertility remains a lifelong mystery.

Key Concepts

1. Male infertility is due to a low sperm count or sperm that cannot swim or are abnormal in structure.
2. Female infertility can be due to an irregular menstrual cycle or blocked uterine tubes. Fibroid tumors, endometriosis, or a misshapen uterus may prevent implantation of a fertilized ovum, and secretions in the vagina and cervix may inactivate or immobilize sperm. Oocytes may fail to release a sperm-attracting biochemical.
3. Early pregnancy loss due to abnormal chromosome number may be mistaken for infertility; this is more common among older women.
4. A variety of medical tests can pinpoint some causes of infertility.

21.3 Assisted Reproductive Technologies

Many people with fertility problems use alternative ways to conceive. Several of the ARTs were developed in nonhuman animals (see the Technology Timeline on page 419). In the United States, about 1 percent of the approximately 4 million births a year are from ARTs, and worldwide ART accounts for about 250,000 births a year.

This section describes types of ARTs. The different procedures can be performed on material from the parents-to-be (“nondonor”) or from donors, and may be “fresh” (collected just prior to the procedure) or “frozen” (preserved in liquid nitrogen).

Donated Sperm—Intrauterine Insemination

The oldest assisted reproductive technology is **intrauterine insemination (IUI)**, in which a doctor places donated sperm into a woman’s cervix or uterus. (It used to be called artificial insemination.) The success rate is 5 to 15% per attempt. The sperm are first washed free of seminal fluid, which can inflame female tissues. A woman might seek IUI if her partner is infertile or carries a gene for an inherited illness that the couple wishes to avoid passing to their child. Women also undergo IUI to be a single parent without having sex.

Technology Timeline

LANDMARKS IN REPRODUCTIVE TECHNOLOGY

	In Nonhuman Animals	In Humans
1782	Intrauterine insemination in dogs	
1790		Pregnancy reported from intrauterine insemination (IUI)
1890s	Birth from embryo transplantation in rabbits	IUI by donor
1949	Cryoprotectant successfully freezes animal sperm	
1951	First calf born after embryo transplantation	
1952	Live calf born after insemination with frozen sperm	
1953		First reported pregnancy after insemination with frozen sperm
1959	Live rabbit offspring produced from <i>in vitro</i> ("test tube") fertilization (IVF)	
1972	Live offspring from frozen mouse embryos	
1976		First reported commercial surrogate motherhood arrangement in the United States
1978	Transplantation of ovaries from one cow to another	Baby born after <i>in vitro</i> fertilization (IVF) in United Kingdom
1980		Baby born after IVF in Australia
1981	Calf born after IVF	Baby born after IVF in United States
1982	Sexing of embryos in rabbits Cattle embryos split to produce genetically identical twins	
1983		Embryo transfer after uterine lavage
1984		Baby born in Australia from frozen and thawed embryo
1985		Baby born after gamete intrafallopian transfer (GIFT) First reported gestational surrogacy arrangement in the United States
1986		Baby born in the United States from frozen and thawed embryo
1989		First preimplantation genetic diagnosis (PGD)
1992		First pregnancies from intracytoplasmic sperm injection (ICSI)
1994	Intracytoplasmic sperm injection (ICSI) in mouse and rabbit	62-year-old woman gives birth from donated oocyte
1995	Sheep cloned from embryo cell nuclei	Babies born following ICSI
1996	Sheep cloned from adult cell nucleus	
1998	Mice cloned from adult cell nuclei	Baby born 7 years after his twin
1999	Cattle cloned from adult cell nuclei	
2000	Pigs cloned from adult cell nuclei	
2001		Sibling born following PGD to treat sister for genetic disease Human preimplantation embryo cloned, survives to 6 cells
2003		3000+ preimplantation genetic diagnoses performed to date
2004	Woman pays \$50,000 to have her cat cloned	First birth from a woman who had ovarian tissue preserved and implanted on an ovary after cancer treatment
2005	Dog cloned	
2009		A woman who had already given birth to six children through IVF has octuplets.

The first documented IUI in humans was done in 1790. For many years, physicians donated sperm, and this became a way for male medical students to earn a few extra dollars. By 1953, sperm could be frozen and stored and IUI became much more commonplace. Today, donated sperm are frozen and stored in sperm banks, which provide the cells to obstetricians who perform the procedure. IUI costs about \$125 to \$615, with higher charges from some facilities for sperm from donors who have professional degrees because those men are paid more for their donations. Additional fees are charged for a more complete medical history of the donor, for photos of the man at different ages, and for participation in a “consent program” in which the donor’s identity is revealed when his offspring turns 18 years old. If ovulation is induced to increase the chances of success of IUI, additional costs may exceed \$3,000.

A couple who chooses IUI can select sperm from a catalog that lists the personal characteristics of donors, such as blood type, hair and eye color, skin color, build, educational level, and interests. One donor profile listed that the man enjoys spear-fishing and wrestling, listens to singer Tori Amos, and loves the film *The Princess Bride*, as if these are inherited traits. Many women selected him because he was a handsome doctor! If a couple desires a child of one sex—such as a daughter to avoid passing on an X-linked disorder—sperm can be separated into fractions enriched for X-bearing or Y-bearing sperm.

Problems can arise in IUI if a donor learns that he has an inherited disease. For example, a man developed cerebellar ataxia (MIM 608029), a movement disorder, years after he donated sperm. Eighteen children conceived using his sperm face a 1 in 2 risk of having inherited the mutant gene. Over-enthusiastic sperm donors can lead to problems. One man, listed in the Fairfax Cryobank as “Donor 401,” earned \$40,000 donating sperm while in law school. He was quite attractive and popular, and forty-five children were conceived with his sperm. When a few of the families he started appeared on a talk show, several other families tuning in were shaken to see so many children who resembled their own. The website <http://www.donorsiblingregistry.com> enables half-siblings who share sperm donor fathers to meet. Thousands have already done so. Each year thousands of newborns trace their beginnings to intrauterine insemination.

A male’s role in reproductive technologies is simpler than a woman’s. A man can be a genetic parent, contributing half of his genetic self in his sperm, but a woman can be both a genetic parent (donating an oocyte) and a gestational parent (donating the uterus). Problems can arise when a second female assists in conception and/or gestation.

A Donated Uterus—Surrogate Motherhood

If a man produces healthy sperm but his partner’s uterus cannot maintain a pregnancy, a surrogate mother may help by being inseminated with the man’s sperm. When the child is born, the surrogate mother gives the baby to the couple. In this variation of the technology, the surrogate is both the genetic and the gestational

mother. Attorneys usually arrange surrogate relationships. The surrogate mother signs a statement signifying her intent to give up the baby. In some U.S. states, and in some nations, she is paid for her 9-month job, but in the United Kingdom compensation is illegal. This is to prevent wealthy couples from taking advantage of women who become surrogates for the money.

A problem with surrogate motherhood is that a woman may not be able to predict her responses to pregnancy and childbirth in a lawyer’s office months earlier. When a surrogate mother changes her mind about giving up the baby, the results are wrenching for all. A prominent early case involved Mary Beth Whitehead, who carried the child of a married man for a fee and then changed her mind about giving up the baby. The courts eventually awarded custody to the father and his wife.

Another type of surrogate mother lends only her uterus, receiving a fertilized ovum conceived from a man and a woman who has healthy ovaries but lacks a functional uterus. This variation is an “embryo transfer to a host uterus.” The gestational-only surrogate mother turns the child over to the biological parents.

In Vitro Fertilization

In *in vitro* fertilization (IVF), which means “fertilization in glass,” sperm and oocyte join in a laboratory dish. Soon after, the embryo that forms is placed in the oocyte donor’s uterus (or another woman’s uterus). If all goes well, it implants into the uterine lining and continues development.

Louise Joy Brown, the first “test-tube baby,” was born in 1978, amid great media attention and sharp criticism. Cartoons depicted “test tube babies,” a great exaggeration of reality. At the time, a prominent bioethicist said that IVF challenged “the idea of humanness and of our human life and the meaning of our embodiment and our relation to ancestors and descendants.” Yet Louise is, despite her unusual beginnings, an ordinary young woman. IVF has since led to the births of more than 4 million children. Today IVF accounts for one in eighty births in the United States.

A woman might undergo IVF if her ovaries and uterus work but her uterine tubes are blocked. Using a laparoscope, which is a lit surgical instrument inserted into the body through a small incision, a physician removes several of the largest oocytes from an ovary and transfers them to a culture dish. If left in the body, only one oocyte would exit the ovary, but in culture, many can mature sufficiently to be fertilized *in vitro*. Chemicals, sperm, and other cell types similar to those in the female reproductive tract are added to the culture. An acidic solution may be applied to the zona pellucida, which is the layer around the egg, to thin it to ease the sperm’s penetration.

Sperm that cannot readily enter the oocyte may be sucked up into a tiny syringe and microinjected into the female cell. This technique, called **intracytoplasmic sperm injection** (ICSI), is more effective than IVF alone and has become standard at some facilities (**figure 21.3**). ICSI is very helpful for men who have low sperm counts or many abnormal sperm, and makes fatherhood possible for men who cannot ejaculate, such as those who have suffered spinal cord injuries. ICSI has been

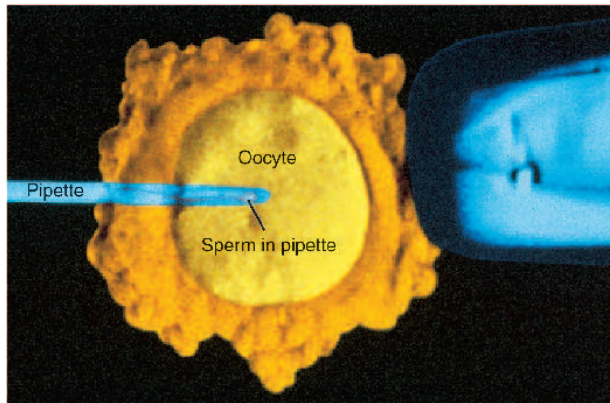


Figure 21.3 ICSI. Intracytoplasmic sperm injection (ICSI) enables some infertile men, men with spinal cord injuries, or men with certain illnesses to become fathers. A single sperm cell is injected into the cytoplasm of an oocyte. This photo is falsely colored.

performed on thousands of men with about a 30 percent success rate. The *Bioethics: Choices for the Future* on page 422 considers a problem with ICSI—transmitting infertility.

Five days after sperm wash over the oocytes in the dish, or are injected into them, one or two blastocysts are transferred to the uterus. If the hormone human chorionic gonadotropin appears in her blood a few days later, and its level rises, she is pregnant.

IVF costs from \$6,500 to \$15,000 per attempt. Children born following IVF have a slight increase in the rate of birth defects (about 8 percent) compared to children conceived naturally. This may reflect medical problems of parents seeking the procedure, dangers of the procedures, the closer scrutiny of IVF pregnancies, and/or effects on imprinting from the few days that the embryo was in culture.

In the past, several embryos were implanted to increase the success rate of IVF, but this led to many multiple births. In many cases, embryos had to be removed to make room for others to survive. To avoid the multiples problem, and because IVF has become more successful as techniques have improved, guidelines now suggest transferring only one embryo.

Embryos resulting from IVF that are not soon implanted in the woman can be frozen in liquid nitrogen (“cryopreserved”) for later use. Cryoprotectant chemicals are used to prevent salts from building up or ice crystals from damaging delicate cell parts. Freezing takes a few hours; thawing about a half hour. The longest an embryo has been frozen, stored, and then successfully revived is 13 years; the “oldest” pregnancy using a frozen embryo occurred 9 years after the freezing!

Gamete and Zygote Intrafallopian Transfer

IVF may fail because of the artificial environment for fertilization. A procedure called GIFT, which stands for **gamete intrafallopian transfer**, improves the setting. (Uterine tubes are also called fallopian tubes). Fertilization is assisted in GIFT, but it occurs in the woman’s body rather than in glassware.

In GIFT, a woman has several of her largest oocytes removed. The man submits a sperm sample, and the most active cells are separated from it. The collected oocytes and sperm are deposited together in the woman’s uterine tube, at a site past any obstruction that might otherwise block fertilization. GIFT is about 22 percent successful, and usually costs less than IVF.

A variation of GIFT is ZIFT, which stands for **zygote intrafallopian transfer**. In this procedure, an IVF ovum is introduced into the woman’s uterine tube. Allowing the fertilized ovum to make its own way to the uterus increases the chance that it will implant. ZIFT is also 22 percent successful.

GIFT and ZIFT are done less frequently than IVF. They often will not work for women who have scarred uterine tubes.

Oocyte Banking and Donation

Oocytes can be stored, as sperm are, but the procedure may create problems. An oocyte contains a large volume of water. Freezing can form ice crystals that damage cell parts. Candidates for preserving oocytes include women who wish to have children later in life and women who will contact toxins or teratogens in the workplace or in chemotherapy.

Oocytes are frozen in liquid nitrogen at -30 to -40 degrees Celsius, when they are at metaphase of the second meiotic division. At this time, the chromosomes are aligned along the spindle, which is sensitive to temperature extremes. If the spindle comes apart as the cell freezes, the oocyte may lose a chromosome, which would devastate development. Another problem with freezing oocytes is retention of a polar body, leading to a diploid oocyte. Only 100 babies have been born using frozen oocytes despite two decades of attempts. The probability of pregnancy using a frozen oocyte with current technology is only about 3 percent.

To avoid the difficulty of freezing oocytes, strips of ovarian tissue can be frozen, stored, thawed, and reimplanted at various sites, such as under the skin of the forearm or abdomen or in the pelvic cavity near the ovaries. The tissue ovulates and the oocytes are collected and fertilized *in vitro*. The first child resulting from fertilization of an oocyte from reimplanted ovarian tissue was born in 2004. The mother, age 25, had been diagnosed with advanced Hodgkin’s lymphoma. The harsh chemotherapy and radiation cured her cancer, but destroyed her ovaries. Beforehand, 5 strips of tissue from her left ovary were frozen. Later, several pieces of ovarian tissue were thawed and implanted in a pocket that surgeons crafted on one of her shriveled ovaries, near the entrance to a uterine tube. Menstrual cycles resumed, and shortly thereafter, the woman became pregnant with her daughter, who is healthy. Freezing ovarian tissue is likely to become routine for cancer patients of childbearing age.

Women can also obtain oocytes from donors, typically younger women. The recipients might not make oocytes or wish to avoid passing on a mutation. Often the donors are undergoing IVF and have “extra” harvested oocytes (see the opening essay for chapter 3 on oocyte donation). The potential father’s sperm and donor’s oocytes are placed in the recipient’s uterus or uterine tube, or fertilization occurs in the laboratory, and a blastocyst is transferred to the woman’s uterus. A program in the U.K. funds IVF for women who cannot afford the procedure if they donate

Bioethics: Choices for the Future

Technology Too Soon? The Case of ICSI

Intracytoplasmic sperm injection (ICSI) has enabled many men with AIDS, paralysis, very low sperm counts, or abnormal sperm to become fathers, without increasing the birth defect rate. But potential problems are emerging, based on the fact that ICSI bypasses what one researcher calls “natural sperm selection barriers.”

ICSI is now commonly used on men who make very few or no sperm by using a needle to extract precursor cells, called spermatids, from the testes. That is, the infertility does not arise until the sperm mature. About 10 percent of infertile men have microdeletions in the Y chromosome and when they use ICSI, they pass on the infertility to their sons.

Bioethicists debate whether it is right to intentionally conceive a male who is genetically destined to be infertile. On the positive side, sons would know of the Y chromosome microdeletions, and perhaps store their spermatids for later use. In the past, these men would not have suspected that they had Y chromosome abnormalities until they had difficulty fathering children. Alternatives to transmitting deletions and other mutations include selecting and using only X-bearing sperm in ICSI, or selecting and implanting only female (XX) fertilized ova. Because of the transmission of Y-linked infertility with ICSI, men undergoing fertility testing and considering the procedure now have their Y chromosomes screened for deletions, and genetic counseling is provided.

Experiments on rhesus monkeys are pinpointing the sources of damage to ICSI embryos that cease developing, which may reveal causes of early pregnancy loss in humans. Findings include:

- Injecting sperm at the site of the polar body on the oocyte can disrupt the meiotic spindle, leading to nondisjunction (an extra or missing chromosome).
- Injected sperm DNA does not always condense properly, also leading to nondisjunction.

- Spermatids that have not yet elongated are often unable to fertilize an oocyte. Culturing them in the laboratory until they mature may help improve the odds of success of IVF.
- Spermatids may not be completely imprinted, leading to problems in gene expression.
- Mitotic cell cycle checkpoints are altered at the first division following ICSI.
- Injected sperm sometimes lack a protein that normally associates with the sex chromosomes. This may explain an elevation in sex chromosome anomalies seen among ICSI children.
- Injected sperm can include surface proteins normally left outside the oocyte, producing unanticipated effects. They may also include mitochondria from the male.

Despite these largely theoretical concerns, ICSI seems to be remarkably safe in humans. Ongoing studies are following the children of ICSI and investigating any associations between health problems in offspring and the cause of subfertility or infertility in the fathers.

Questions for Discussion

1. Does helping a person to see with glasses have the same consequences to the next generation as ICSI for a man with chromosome-based infertility?
2. What does ICSI offer compared to adoption?
3. Is it ethical for a man with a Y chromosome deletion to use ICSI to have a son?
4. Would it be ethical to conduct the types of experiments done with ICSI in monkeys on humans?

their “extras.” The success of using oocytes from younger women confirms that it is the oocyte that age affects, and not the uterine lining. The live birth rate per cycle for older women using their own oocytes in IVF is 28 percent. For older women using oocytes donated by younger women, the rate is about 50 percent.

Embryo adoption is a variation on oocyte donation. A woman with malfunctioning ovaries but a healthy uterus carries an embryo that results when her partner’s sperm is used in intrauterine insemination of a woman who produces healthy oocytes. If the woman conceives, the embryo is gently flushed out of her uterus a week later and inserted through the cervix and into the uterus of the woman with malfunctioning ovaries. The child is genetically that of the man and the woman who carries it for the first week, but is born from the woman who cannot produce healthy oocytes. “Embryo adoption” also describes use of IVF “leftovers.”

In another technology, cytoplasmic donation, older women have their oocytes injected with cytoplasm from the oocytes of younger women to “rejuvenate” the cells. Although resulting children conceived through IVF appear to be healthy, they

are being monitored for a potential problem—heteroplasmy, or two sources of mitochondria in one cell. Researchers do not yet know the health consequences of having mitochondria from the donor cytoplasm plus mitochondria from the recipient’s oocyte. These conceptions also have an elevated incidence of XO syndrome, which often causes spontaneous abortion.

Because oocytes are harder to obtain than sperm, oocyte donation technology has lagged behind that of sperm banks, but is catching up. One IVF facility that has run a donor oocyte program since 1988 has a patient brochure that describes 120 oocyte donors of various ethnic backgrounds, like a catalog of sperm donors. The oocyte donors are young and have undergone extensive medical and genetic tests. Recipients may be up to 55 years of age.

Preimplantation Genetic Diagnosis

Prenatal diagnostic tests such as amniocentesis and chorionic villus sampling can be used in pregnancies achieved with assisted reproductive technologies. A test called **preimplantation genetic**

diagnosis (PGD) detects genetic and chromosomal abnormalities *before* pregnancy starts. The couple selects a very early “preimplantation” embryo, so-called because it would not normally have arrived at the uterus for implantation. The selected embryo has not inherited a specific detectable genetic condition. This technology was used to select Adam Nash, whose umbilical cord stem cells cured his sister’s Fanconi anemia (see figure 21.1). PGD has about a 29 percent success rate.

PGD is possible because one cell, or blastomere, can be removed for testing from an eight-celled embryo and the remaining seven cells can complete development normally in a uterus. Before the embryo is implanted into the woman the single cell is karyotyped, or its DNA amplified and probed for genes that the parents carry. Embryos that pass the chromosomal check-up are selected to complete development or are stored. At first, researchers implanted the remaining seven cells, but letting the embryo continue developing in the dish until day 5, when it is 80 to 120 cells, is more successful. Obtaining the cell to be tested is called “blastomere biopsy” (**figure 21.4**). Accuracy is about 97 percent. Errors are generally due to mosaics—when a somatic mutation occurs in a blastomere—or during amplification of blastomere DNA.

The first children who had PGD were born in 1989. In these first cases, probes for Y chromosome-specific DNA sequences were used to select females, who could not inherit X-linked conditions their mothers carried. The alternative to PGD would have been to face the 25 percent chance of conceiving an affected male.

In March 1992, the first child was born who underwent PGD to avoid a specific inherited disease. Chloe O’Brien was checked as an eight-celled preimplantation embryo to see if

she had escaped the cystic fibrosis that affected her brother. Since then, PGD has helped to select thousands of children free of several dozen types of inherited illnesses. It has been used for the better-known single-gene disorders as well as for rare ones.

Today PGD is increasingly being used to screen early embryos derived from IVF for normal chromosome number before implanting them into women. This should increase the chances of successful live births, but in the first large trial, PGD actually lowered the birth rate—perhaps the intervention harms the embryos. In the Netherlands, researchers examined all of the cells of several preimplantation human embryos. They found that, within an embryo, some cells can have normal chromosomes and others not. Therefore, the assumption underlying PGD, that a sampled cell represents them all, may need to be re-examined.

Like many technologies, PGD can introduce a bioethical “slippery slope” when it is used other than to ensure that a child is free of a certain disease, such as for gender selection. A couple with five sons might, for example, use PGD to select a daughter. But this use of technology might just be a high-tech version of age-old human nature, according to one physician who performs PGD. “From the dawn of time, people have tried to control the sex of offspring, whether that means making love with one partner wearing army boots, or using a fluorescence-activated cell sorter to separate X- and Y-bearing sperm. PGD represents a quantum leap in that ability—all you have to do is read the X and Y chromosome paints,” he says.

While PGD used solely for family planning is certainly more civilized than placing baby girls outside the gates of ancient cities to perish, the American Society for Reproductive Medicine

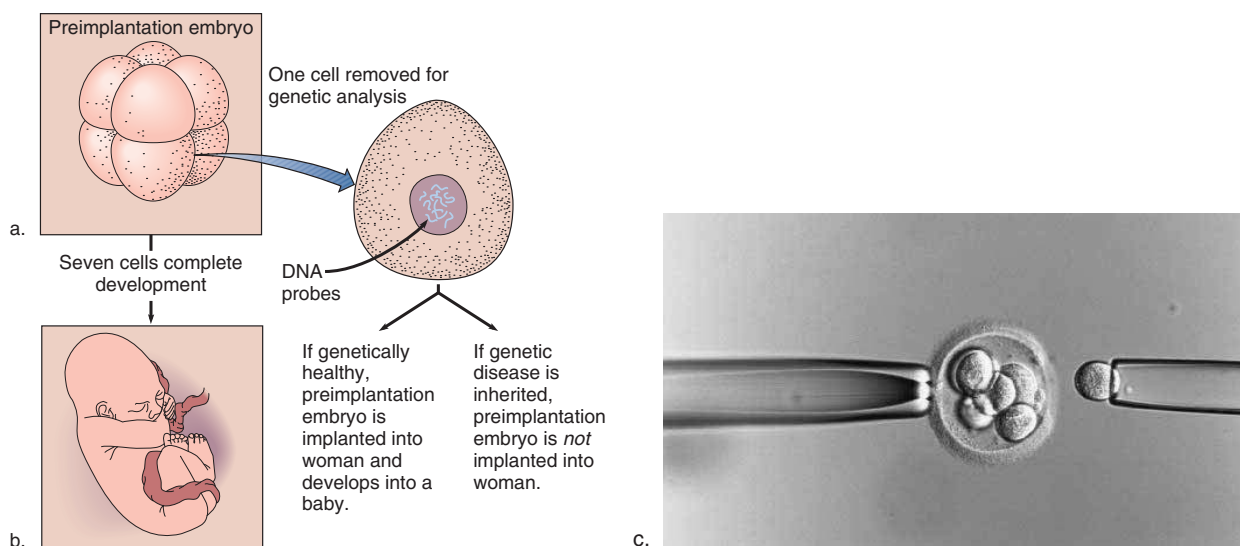


Figure 21.4 Preimplantation genetic diagnosis (PGD) probes disease-causing genes or chromosome aberrations in an 8-celled preimplantation embryo. (a) A single cell is separated and tested to see if it contains a disease-causing genotype or chromosome abnormality. (b) If it doesn’t, the remaining seven cells divide a few more times and are transferred to the oocyte donor to complete development. (c) This preimplantation embryo is held still by suction applied on the left. On the right, a pipette draws up a single blastomere. *In vitro* fertilization took place 45 hours previously.

endorses the use of PGD for sex selection only to avoid passing on an X-linked disease. Yet even PGD to avoid disease can be controversial. In the United Kingdom, where the government regulates reproductive technology, inherited cancer susceptibility is an approved reason to have PGD. However, bioethicists point out that these cancers do not begin until adulthood, the susceptibility is incompletely penetrant (not everyone who inherits the disease-associated genotype will actually develop cancer), and the cancer may be treatable.

Table 21.1 summarizes the assisted reproductive technologies.

Key Concepts

1. Intrauterine insemination places donor sperm in a woman's reproductive tract.
2. A genetic and gestational surrogate mother is intrauterinely inseminated, becomes pregnant, then gives the baby to the father and his partner. A gestational surrogate mother gestates a baby conceived *in vitro* with gametes from a man and a woman who cannot carry a fetus.
3. In IVF, sperm and oocyte unite outside the body, and the resulting embryo is transferred to the uterus. Early embryos can also be frozen and used later.
4. In GIFT, sperm and oocytes are placed in a uterine tube past a blockage. In ZIFT, an IVF embryo is placed in a uterine tube.
5. In embryo adoption, a woman who has had intrauterine insemination has an early embryo washed out of her uterus and transferred to a woman who lacks oocytes.
6. PGD removes cells from early embryos and screens them for genetic or chromosomal abnormalities.

21.4 Extra Embryos

Sometimes assisted reproductive technologies leave “extra” oocytes, fertilized ova, or very early embryos. **Table 21.2** lists the possible fates of this biological material.

In the United States, nearly half a million embryos derived from IVF sit in freezers; some have been there for years. Most couples who donate embryos to others do so anonymously, with no intention of learning how their genetic offspring are raised. Scott and Glenda Lyons chose a different path when they learned that their attempt at IVF had yielded too many embryos.

In 2001, two of Glenda's eighteen embryos were transferred to her uterus, and developed into twins Samantha and Mitchell. Through a website where couples chat about fertility issues, Scott and Glenda met and selected Bruce and Susan Lindeman to receive fourteen remaining embryos. This second couple had tried IVF three times, with no luck. The Lyoneses frozen embryos were shipped cross-country to a clinic where two were implanted in Susan's uterus. In July 2003, Chase and Jack Lindeman were born—genetic siblings of Samantha and Mitchell Lyones. But there were still embryos left. The Lyons allowed the Lindemans to send twelve embryos to a third couple, who used two to have twin daughters in August 2004. They are biological siblings of Samantha and Mitchell Lyons and Chase and Jack Lindeman (**figure 21.5**). Like the Lyoneses a woman in California who had eight fertilized ova left over after using six to produce her six young children, also did not want to destroy them or continue to store them. Nadya Suleman was implanted with them, and in early 2009 gave birth to octuplets!

Donating fertilized ova and embryos for use in research is another alternative to disposing of them. The results of these experiments sometimes challenge long-held ideas, indicating that we still have much to learn about early human prenatal development. This

Table 21.1 Some Assisted Reproductive Technologies

Technology	Procedures	Success/Cycle	Cost/Cycle
GIFT	Deposits collected oocytes and sperm in uterine tube.	27%	\$8,000–\$10,000
IVF	Mixes sperm and oocytes in a laboratory dish, with chemicals to simulate intrauterine environment to encourage fertilization.	29%	\$6,500–\$15,000
IUI	Places or injects washed sperm into the cervix or uterus.	5–25%	\$125
ICSI	Injects immature or rare sperm into oocyte, before IVF.	28%	\$10,000–\$17,000
Oocyte freezing	Oocytes retrieved and frozen in liquid nitrogen.	3%	\$8,000
Ovulation induction	Drugs control timing of ovulation in order to perform a particular procedure.	28%	\$3,000
PGD	Searches for specific mutant allele in sampled cell of 8-celled embryo. Its absence indicates remaining 7-celled embryo can be nurtured and implanted in woman, and child will be free of genetic condition.	29%	\$8,000–\$15,000
Surrogate mother	Woman carries a pregnancy for a woman who cannot become or stay pregnant.		\$10,000
ZIFT	Places IVF ovum in uterine tube.	29%	\$8,000–\$13,000

was the case for a study from Royal Victoria Hospital in Montreal. Researchers examined the chromosomes of sperm from a man with XXY syndrome. Many of the sperm would be expected to have an extra X chromosome, due to nondisjunction (see figure 13.12), which could lead to a preponderance of XXX and XXY offspring. Surprisingly, only 3.9 percent of the man's *sperm* had extra

chromosomes, but five out of ten of his spare *embryos* had an abnormal X, Y, or chromosome 18. That is, even though most of the man's sperm were normal, his embryos weren't. The source of reproductive problems in XXY syndrome, therefore, might not be in the sperm, but in early embryos—a finding that was previously unknown and not expected, and was only learned because of observing early human embryos.

In another study, Australian researchers followed the fates of single blastomeres that had too many or too few chromosomes. They wanted to see whether the abnormal cells preferentially ended up in the inner cell mass, which develops into the embryo, or the trophoblast, which becomes extra-embryonic membranes. The study showed that cells with extra or missing chromosomes become part of the inner cell mass much more frequently than expected by chance. This finding indicates that the ability of a blastomere sampled for PGD to predict health may depend on whether it is fated to be part of the inner cell mass.

Using fertilized ova or embryos designated for discard in research is controversial. Without regulations on privately funded research, ethically questionable experiments can happen. For example, researchers reported at a conference that they had mixed human cells from male embryos with cells from female embryos, to see if the normal male cells could "save" the female cells with a mutation. Sex was chosen as a marker because the Y chromosome is easy to detect. But the idea of human embryos with mixed sex parts caused a public outcry.

In vitro fertilized ova not chosen for reproduction have been the major source of donations for research. However, an experimental ART called **polar body biopsy** may reduce this supply by increasing the success of IVF. Polar body biopsy is based on Mendel's first law, the segregation of alleles. In the technique, if a polar body resulting from the first meiotic division in a woman who is a carrier of an X-linked disorder has the mutant allele, then the oocyte to which it clings lacks that allele. Oocytes that pass this test can be fertilized *in vitro* and the resulting embryo implanted. Polar body biopsy is possible because the polar body is attached to the much larger oocyte. A large pipette is used to hold the two cells in place, and a smaller pipette is used to separate the polar body. Then, DNA probes and FISH are used to look at genes and chromosomes in the polar body and infer the genotype of the oocyte (figure 21.6). Polar body biopsy followed by PGD is quite

Table 21.2 Fates of Frozen Embryos

1. Store indefinitely.
2. Store and destroy after a set time.
3. Donate for embryonic stem cell derivation and research.
4. Thaw later for use by biological parents.
5. Thaw later for use by other parents.
6. Discard.

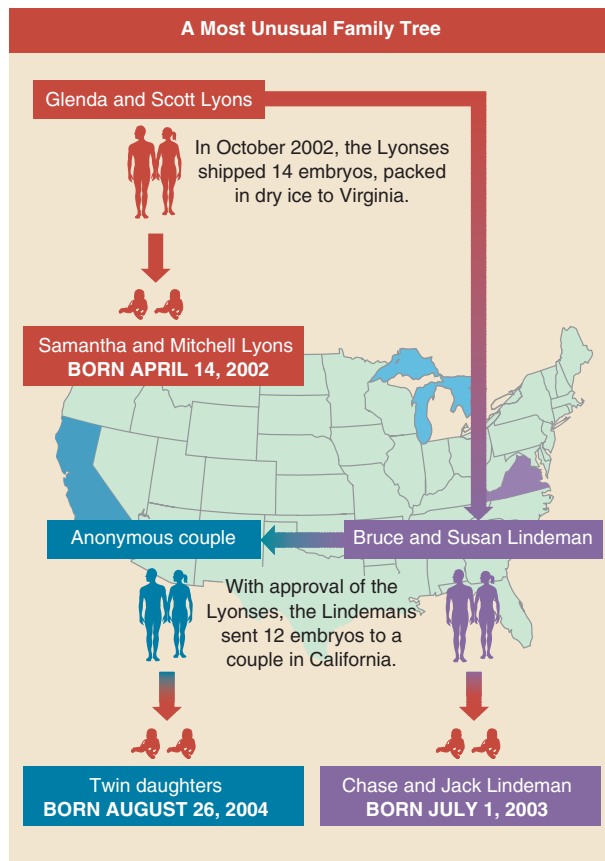


Figure 21.5 Using extra embryos. Six children resulted from Glenda and Scott Lyons' embryos. The Lyonses had a boy and a girl, then donated embryos to the Lindemans, who had twin boys. Finally, a couple in California used the Lyonses embryos to have twin daughters. The Lyonses and Lindemans have become friends.



Figure 21.6 Polar body biopsy. The fact that an oocyte shares a woman's divided genetic material with a much smaller polar body allows the screening of oocytes for use in IVF.

effective in avoiding conceptions with chromosome abnormalities or certain single-gene disorders.

ARTs introduce ownership and parentage issues (table 21.3). Another controversy is that human genome information is providing more traits to track and perhaps control in coming generations. When we can routinely scan the human genome in gametes, fertilized ova, or early embryos, who will decide which traits are worth living with, and which aren't?

ARTs operate on molecules and cells, but with repercussions for individuals and families. Ultimately, if it becomes widespread enough, these interventions into reproduction may affect the gene pool. Let us hope that regulations will evolve along with the technologies to assure that they are applied sensibly and humanely.

Key Concepts

1. IVF produces extra fertilized ova and early embryos that may be used, frozen, donated, discarded, or used in research.
2. Embryos in research add to our knowledge of early human development.
3. Polar body biopsy enables physicians to select out defective oocytes.

Table 21.3 Assisted Reproductive Disasters

1. A physician in California used his own sperm to perform intrauterine insemination on 15 patients, telling them that he had used sperm from anonymous donors.
2. A plane crash killed the wealthy parents of two early embryos stored at -320°F (-195°C) in a hospital in Melbourne, Australia. Adult children of the couple were asked to share their estate with two 8-celled siblings-to-be.
3. Several couples in Chicago planning to marry discovered that they were half-siblings. Their mothers had been inseminated with sperm from the same donor.
4. Two Rhode Island couples sued a fertility clinic for misplacing several embryos.
5. Several couples in California sued a fertility clinic for implanting their oocytes or embryos in other women without donor consent. One woman requested partial custody of the resulting children if her oocytes were taken, and full custody if her embryos were used, even though the children were of school age and she had never met them.
6. A man sued his ex-wife for possession of their frozen fertilized ova. He won, and donated them for research. She had wanted to be pregnant.
7. The night before *in vitro* fertilized embryos were to be implanted in a 40-year-old woman's uterus after she and her husband had spent four years trying to conceive, the man changed his mind, and wanted the embryos destroyed. Their embryo custody battle is before the Supreme Court.

Summary

21.1 Savior Siblings and More

1. **Assisted reproductive technologies** replace what is missing in reproduction.

21.2 Infertility and Subfertility

2. **Infertility** is the inability to conceive a child after a year of unprotected intercourse. Subfertile individuals or couples manufacture gametes, but take longer than usual to conceive.
3. Causes of infertility in the male include low sperm count, a malfunctioning immune system, a varicose vein in the scrotum, structural sperm defects, drug exposure, vasectomy reversal, and abnormal hormone levels. Mutation may impair fertility.
4. Female infertility can be caused by absent or irregular ovulation, blocked uterine tubes, an inhospitable or malshaped uterus, antisperm secretions, or lack of sperm-attracting biochemicals. Early pregnancy loss due to abnormal chromosome number is more common in older women and may appear to be infertility.

21.3 Assisted Reproductive Technologies

5. In **intrauterine insemination**, donor sperm are introduced into a woman's reproductive tract in a clinical setting.
6. A gestational and genetic surrogate mother provides her oocyte. Then intrauterine insemination is performed with

sperm from a man whose partner cannot conceive or carry a fetus. The surrogate also provides her uterus for nine months. A gestational surrogate mother receives an ***in vitro* fertilized ovum** that belongs genetically to the couple who ask her to carry it.

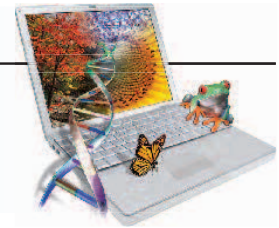
7. In IVF, oocytes and sperm meet in a dish, fertilized ova divide a few times, and embryos are placed in the woman's body, circumventing blocked tubes or malfunctioning sperm. **Intracytoplasmic sperm injection** introduces immature or nonmotile sperm into oocytes.
8. Embryos can be frozen and thawed and then complete development when placed in a woman's uterus.
9. **GIFT** introduces oocytes and sperm into a uterine tube past a blockage; fertilization occurs in the woman's body. **ZIFT** places an early embryo in a uterine tube.
10. Oocytes can be frozen and stored. In embryo adoption, a woman undergoes intrauterine insemination. A week later, the embryo is washed out of her uterus and introduced into the reproductive tract of the woman whose partner donated the sperm.
11. Seven-celled embryos can develop normally if a blastomere is removed at the eight-cell stage and cleared for abnormal chromosomes or genes. This is **preimplantation genetic diagnosis**.

21.4 Extra Embryos

- Extra fertilized ova and early embryos generated in IVF are used, donated to couples, stored, donated for research, or discarded. They enable researchers to study aspects of early human development that they could not investigate in other ways.
- Polar body biopsy** enables physicians to perform genetic tests on polar bodies and to infer the genotype of the accompanying oocyte.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

- Which assisted reproductive technologies might help the following couples? (More than one answer may fit some situations.)
 - A woman is born without a uterus, but manufactures healthy oocytes.
 - A man has cancer treatments that damage his sperm.
 - A woman undergoes a genetic test that reveals she will develop Huntington disease. She wants to have a child, but does not want to pass on this presently untreatable illness.
 - Two women wish to have and raise a child together.
 - A man and woman are each carriers of sickle cell disease. They do not want to have an affected child, but they also do not want to terminate a pregnancy.
 - A woman's uterine tubes are scarred and blocked, so an oocyte cannot reach the uterus.
 - A young woman must undergo abdominal radiation to treat ovarian cancer, but wishes to have a child.
- Why are men typically tested for infertility before women?
- A man reads his medical chart and discovers that the results of his sperm analysis indicate that 22 percent of his sperm are shaped abnormally. He wonders why the physician said he had normal fertility if so many sperm are abnormally shaped. Has the doctor made an error?
- Cite a situation in which both man and woman contribute to subfertility.
- How does ZIFT differ from GIFT? How does it differ from IVF?
- A Tennessee lower court, in ruling on the fate of seven frozen embryos in a divorce case, called them "children *in vitro*." In what sense is this label incorrect?
- Explain how preimplantation genetic diagnosis is similar to and different from CVS and amniocentesis, described in chapter 13.
- What are some of the causes of infertility among older women?
- How do each of the following assisted reproductive technologies deviate from the normal biological process?
 - in vitro* fertilization
 - GIFT
 - embryo adoption
 - gestational-only surrogacy
 - intrauterine insemination
 - cytoplasmic donation
- When Louise Joy Brown was born in 1978, many people were horrified and several government officials called for banning IVF. Today, more than 4 million people have been conceived using the technology. Are there any experimental treatments or technologies today that people are wary of that might one day become routine?
- Explain how PGD works.

Applied Questions

- The San Antonio Embryo Bank is the first such facility in the United States. It offers IVF leftover embryos, which would otherwise remain in the deep freeze or be discarded, to people wanting to have children, for \$2,500 each. The bank circumvents bioethical concerns by claiming that it sells a service, not an embryo. People in favor of the bank claim that purchasing an embryo is not different from paying for sperm or eggs, or an adopted child. Those who object to the bank claim that it makes an embryo a commodity.
 - Do you think that an embryo bank is a good idea, or is it unethical?
 - Whose rights are involved in the operation of the embryo bank?
 - Who should be liable if a child that develops from the embryo has a medical problem?
 - Is the bank elitist because the cost is so high?
- A newspaper columnist wrote that frozen human embryos are "microscopic Americans." Former president George W. Bush called them "unique and genetically complete, like every other human being." A stem cell researcher referred to embryonic stem cells as "like any other cell in an adult, no different from the skin cells you rub off with a towel after a shower."

- a. What is your opinion of the status of an eight-cell human embryo?
 - b. What is your opinion of the status of a cell that descends from a cell removed from an eight-cell human embryo? Does the status depend upon the fate of the rest of the embryo?
 - c. Do you think that there is any harm in an influential person, such as a journalist, politician, or researcher, stating the status of an embryo?
 - d. How might the following individuals respond to or feel about these definitions?
 - i. A woman who ended a pregnancy, for whatever reason
 - ii. A couple who have had a spontaneous abortion
 - iii. A stem cell researcher
 - iv. A person with a disease that one day may be treated with stem cells
 - v. A couple who have tried to conceive for a decade
3. At the same time that 62- and 63-year-old women gave birth, actors Tony Randall and Anthony Quinn became fathers at ages 77 and 78—and didn't receive nearly as much criticism as the women. Do you think this is an unfair double standard, or a fair criticism based on valid biological information?
 4. Many people spend thousands of dollars pursuing pregnancy. What is an alternative solution to their quest for parenthood?
 5. An Oregon man anonymously donated sperm that were used to conceive a child. The man later claimed, and won, rights to visit his child. Is this situation for the man more analogous to a genetic and gestational surrogate mother, or an oocyte donor who wishes to see the child she helped to bring into existence?
 6. Big Tom is a bull with valuable genetic traits. His sperm are used to conceive 1,000 calves. Mist, a dairy cow with exceptional milk output, has many oocytes removed, fertilized *in vitro*, and implanted into surrogate mothers. With their help, Mist becomes the genetic mother of 100 calves—far more than she could give birth to naturally. Which two reproductive technologies performed on humans are based on these two agricultural examples?
 7. State who the genetic parents are and who the gestational mother is in each of the following cases:
 - a. A man was exposed to unknown burning chemicals and received several vaccines during the first Gulf war, and abused drugs for several years before and after that. Now he wants to become a father, but he is concerned that past exposures to toxins have damaged his sperm. His wife undergoes intrauterine insemination with sperm from the husband's brother, who has led a calmer and healthier life.
 - b. A 26-year-old woman has her uterus removed because of cancer. However, her ovaries are intact and her oocytes are healthy. She has oocytes removed and fertilized *in vitro* with her husband's sperm. Two resulting embryos are implanted into the uterus of the woman's best friend.
 - c. Max and Tina had a child by IVF in 1986. At that time, they had three extra embryos frozen. Two are thawed years later and implanted into the uterus of Tina's sister, Karen. Karen's uterus is healthy, but she has ovarian cysts that often prevent her from ovulating.
 - d. Forty-year-old Christensen von Wormer wanted children, but not a partner. He donated sperm, which were used for intrauterine insemination of an Indiana mother of one. The woman carried the resulting fetus to term for a fee. On September 5, 1990, von Wormer held his newborn daughter, Kelsey, for the first time.
 - e. Two men who live together want to raise a child. They go to a fertility clinic, have their sperm collected, mixed, and used to inseminate a friend. Nine months later she turns the baby over to them.
8. Delaying childbirth until after age 35 is associated with certain physical risks, yet an older woman is often more mature and financially secure. Many women delay childbirth so that they can establish careers. Can you suggest societal changes, perhaps using a reproductive technology, that would allow women to more easily have children and careers?
 9. An IVF attempt yields twelve more embryos than the couple who conceived them can use. What could they do with the extras?
 10. What do you think children born of an assisted reproductive technology should be told about their origins?
 11. Wealthy couples could hire poor women as surrogates or oocyte donors simply because the adoptive mother does not want to be pregnant. Would you object to this practice? Why or why not?
 12. An IVF program in India offers preimplantation genetic diagnosis to help couples who already have a daughter to conceive a son. The reasoning is that because having a male heir is of such great importance in this society, offering PGD can enable couples to avoid aborting second and subsequent female pregnancies. Do you agree or disagree that PGD should be used for sex selection in this sociological context?

Web Activities

13. The United States government bans use of federal funds to create human embryos for research purposes, but does not regulate the human reproductive technology field at all. Consult websites to learn about regulations of stem cell research and ARTs in other nations.
14. Go to the Centers for Disease Control and Prevention website. Click on ART Trends, and use the information to answer the following questions.
 - a. Since 1996, to what extent has the use of ARTs in the U.S. increased?
 - b. Which is more successfully implanted into an infertile woman's uterus, a fresh or frozen donor oocyte?
 - c. Which is more successfully implanted into an infertile woman's uterus, a donated oocyte or one of her own?
 - d. What are two factors that could complicate data collection on ART success rates?

15. A company called Extend Fertility provides oocyte freezing services, telling women to “set your own biological clock.” The home page states, “Today’s women lead rich and busy lives—obtaining advanced degrees, pursuing successful careers, and taking better care of ourselves. As a result of this progress, many of us choose to have children later than our mothers did.”
- Look at the website. Discuss how it might be viewed by the following individuals:
- A 73-year-old father of a healthy baby
 - A 26-year-old woman, married with no children but who wants them, facing six months of chemotherapy
 - An orphaned 10-year-old in Thailand
 - A healthy 28-year-old woman in the United States who wants to earn degrees in medicine, law, and business before becoming pregnant
 - A young mother in Mexico who is giving her son up for adoption because she cannot afford to raise him
16. Read the posts on surrogatemother.com and describe a match between a potential surrogate and a couple or individual wishing to use her services.

Case Studies and Research Results

17. Doola is 32 years old and is trying to decide if she and her husband are ready for parenthood when she learns that her 48-year-old mother has Alzheimer disease. The mother’s physician tells Doola that because of the early onset, the Alzheimer’s could be inherited through a susceptibility gene. Doola is tested and indeed has the same dominant allele. She wants to have a child right away, so that she can enjoy many years as a mother. Her husband David feels that it wouldn’t be fair to have a child knowing that Alzheimer disease likely lies in Doola’s future.
- Who do you agree with, and why?
 - David is also concerned that Doola could pass on her Alzheimer gene variant to a child. What technology might help them avoid this?
 - Is Doola correct in assuming that she is destined to develop Alzheimer disease?
18. Natallie Evans had to have her ovaries removed at a young age because they were precancerous, so she and her partner had IVF and froze their embryos for use at a later time. Under British law, both partners must consent for the continued storage of frozen embryos. Evans and her partner split, and he revoked his consent. She sued for the right to use the embryos. She told the court, “I am pleased to have the opportunity to ask the court to save my embryos and let me use them to have the child I so desperately want.”
- What information should the court consider in deciding this case? Whose rights do you think should be paramount?
19. Colleen and Ellen were partners who had twin daughters using ART. Colleen was in her forties when they decided to have children, and she had large uterine fibroids, so she could not carry a pregnancy. They selected sperm from a sperm bank, which was used in intrauterine insemination on Ellen, but she did not conceive after several attempts. Next Ellen tried IVF, to no avail. Finally, the women decided to combine their contributions. Colleen had oocytes collected and fertilized and implanted in Ellen’s uterus. The women signed legal documents declaring Ellen the sole parent, with plans to consider a more shared arrangement 5 years later. At the clinic, Colleen signed a form that waived her rights to her oocytes or children resulting from their being fertilized. She would later claim that she signed it because she thought she had to for the procedure to be done. Ellen’s name was recorded on the birth certificates of their twin daughters. Although both women were active parents, their relationship deteriorated, and when the girls were 6, Ellen moved them across the country and kept Colleen from visiting them. The case ended up in the courts, which ruled that Ellen is the mother. The Marin County Superior Court declared Colleen’s relationship with the twins “legally irrelevant.” Colleen has petitioned the Supreme Court of California to consider the case.
- Who is the genetic mother and who is the gestational mother?
 - Do you agree with the Marin County Superior Court’s decision that a genetic relationship is “legally irrelevant?”
 - What information would you need to decide whether Colleen was being discriminated against because she is gay?

PCR #	Size of DNA	Primer #	Comments
# 1	$843 - 372 = 467$	47, 82	HR A-100
# 2	$644 - 125 = 519$	81, 50	HR B-100
# 3	$2421 - 170 = 2251$	57, 55	HR C-100
# 4	$732 - 372 = 360$	53, 55	HR D-100
# 5	$2488 - 170 = 2318$	55, 55	HR E-100
# 6	$72 - 1576 = 516$	57, 55	HR F-100
# 7	$2423 - 170 = 2253$	57, 55	HR G-100
# 8	$2853 - 170 = 2683$	61, 82	HC 12
# 9	$2331 - 170 = 2161$	53, 55	HR H-100

How many genes do we have? Knowing may not be useful.

CHAPTER

22

Genomics

20,000 Genomes and Counting

Genome researchers like numbers. They spent much of the 1990s in anticipation of deciphering the sequence of the 3.2 billion bits of information that comprise a human genome. Shortly before that was accomplished—ahead of schedule—many minds turned instead to discovering the points where human genomes vary. Meanwhile, as DNA sequencing technology improved, the feasibility of one day routinely sequencing people’s genomes grew.

As is true with acquiring new gadgets and gizmos, the first few people to have their genomes sequenced were celebrities and/or millionaires. Craig Venter, the researcher who was first to the finish line in sequencing the human genome in 2001, was also the first to have his genome sequence published, in late 2007. Six months later came publication of the genome sequence of James Watson, co-discoverer of DNA’s structure. In the meantime, ten prominent people joined the Personal Genome Project (“PGP-10”) led by Harvard University geneticist George Church. Long before their genomes were sequenced, the participants talked freely to the media about some of their traits. For example, psychologist Steven Pinker, aka #8, graced the cover of the *New York Times* magazine, but the article explored only a few traits.

Soon after a multinational consortium announced the “1,000 Genomes Project” to establish a catalog of many genomes for the sake of comparison, another group announced a goal of sequencing 20,000 genomes by 2010, assuming the cost will plummet. Those interested in learning about only gene variants that affect health risks can send a saliva sample to the Coriell Personalized Medicine Collaborative to

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participate—free of charge—in a research study of those risks. Only results with “actionable” consequences are reported to individuals, but all of the data will help to provide a view of human genomic variation.

What will all of this genetic information ultimately mean? Will physicians consult strings of A, C, T, and G to determine how best to treat their patients, or will signs, symptoms, family history, and a patient’s observations turn out to be more valuable types of information?

Only time will tell. The idea of knowing our genes makes theoretical sense, but it must be validated with improved clinical outcomes.

22.1 From Genetics To Genomics

Genetics is a young science, genomics younger still. As one field has evolved into another, milestones have come at oddly regular intervals. A century after Gregor Mendel announced and published his findings, the genetic code was deciphered; a century after his laws were rediscovered, the human genome was sequenced.

The term *genome* was coined in 1920 by geneticist H. Winkler. A hybrid of “gene” and “chromosome,” genome then denoted a complete set of chromosomes and its genes. The modern definition refers to all the DNA in a haploid set of chromosomes. The term *genomics*, credited to T. H. Roderick in 1986, indicates the study of genomes. Thoughts of sequencing genomes echoed through much of the twentieth century, as researchers defined and described the units of inheritance from various perspectives.

Beginnings in Linkage Studies

Sequencing the human genome unofficially began in the 1980s with deciphering signposts and developing shortcuts. Many of the initial steps and tools grew from existing technology. Linkage maps from the 1950s and later and many family studies that associated chromosomal aberrations with syndromes enabled researchers to assign some genes to their chromosomes. Then automated DNA sequencing took genetic analysis to a new level—information.

The evolution of increasingly detailed genetic maps is similar to zooming in on a geographical

satellite map (**figure 22.1**). A cytogenetic map is like a map of California within a map of the United States, highlighting only the largest cities. A linkage map is like a map that depicts the smaller cities and large towns, and a physical map is similar to a geographical map indicating all towns. Finally, a sequence map is the equivalent of a Google map showing specific buildings.

Before the human genome project, researchers took a gene-by-gene approach, matching single genes to specific diseases using an approach called **positional cloning**. The technique began with examining a particular phenotype corresponding to a Mendelian disorder in large families, then identifying parts of the genome that only affected relatives shared. Such sequences served as markers of the presumably tightly linked disease-causing gene, and are still used today (see table 5.3).

In the 1980s and 1990s, positional cloning experiments yielded a slow but steady stream of discoveries of genes that cause such diseases as Duchenne muscular dystrophy, cystic fibrosis, and Huntington disease (**Reading 22.1**, page 432).

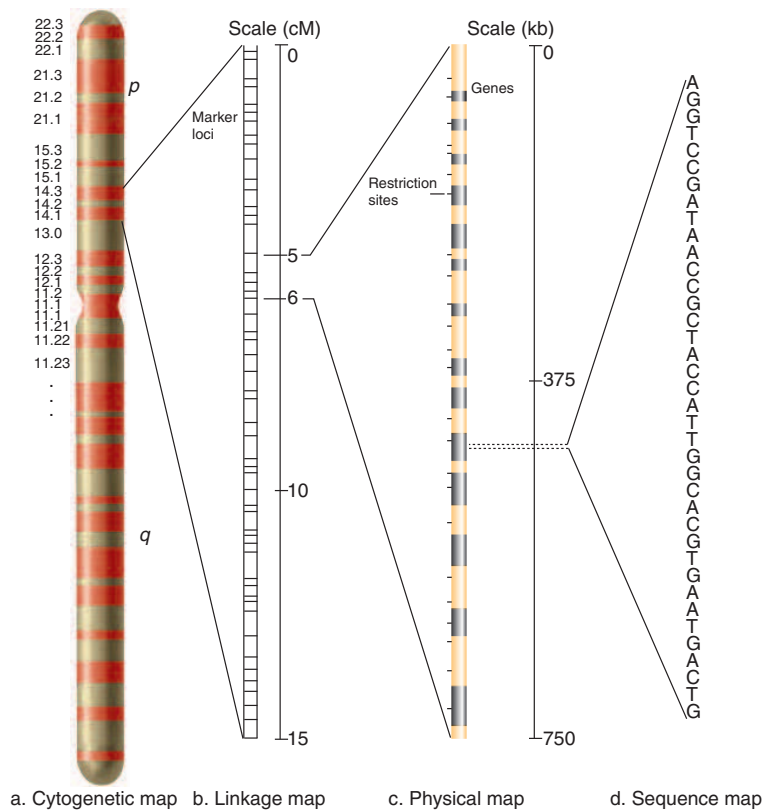


Figure 22.1 Different levels of genetic maps are like zooming up the magnification on a geographical map. (a) A cytogenetic map, based on associations between chromosome aberrations and syndromes, can distinguish DNA sequences that are at least 5,000 kilobases (kb) apart. (b) A linkage map derived from recombination data distinguishes genes hundreds of kb apart. (c) A physical map constructed from overlapped DNA pieces distinguishes genes tens of kb apart. (d) A sequence map depicts the order of nucleotide bases.

Reading 22.1

Discovering the Huntington Disease Gene Took a Decade

Today, finding and sequencing a gene can be done in a day. The story of the discovery of the mutation that causes Huntington disease (HD) illustrates the incredibly slow approach to gene discovery before the human genome sequence was known.

The search for the HD gene began with a large family in a remote village on the shores of Lake Maracaibo, Venezuela. Seven generations ago, in the 1800s, a local woman married a visiting Portuguese sailor who, according to folklore, walked as if intoxicated. Like many couples in the poor fishing village, the woman and her sailor had many children. Some grew up to walk in the same peculiar way as their father. Of their nearly 5,000 descendants, more than 250 living today have HD. This extended family presented a natural experiment to geneticists, and they began by drafting a huge pedigree that eventually depicted more than 10,000 individuals and stretched to more than 100 feet in width (**figure 1**). The Venezuelan family was large enough for researchers to detect a telltale DNA sequence unique to those who had, or would develop, HD—a genetic marker.

In 1981, Columbia University psychologist Nancy Wexler, whose mother would eventually die of HD, began annual visits to Lake Maracaibo. The people lived in huts perched on stilts, as their ancestors did. Wexler traded candy and blue jeans for blood samples and skin biopsies. Meanwhile, investigators at Massachusetts General Hospital were sampling tissue from an Iowa family of forty-one, in which twenty-one individuals had HD. They extracted DNA from the samples, cut it with restriction enzymes, and tested the fragments with a set of labeled DNA probes, looking for one that bound only the DNA of sick people. This would reveal a RFLP (see section 14.4) unique to people with the disease. The team added the Venezuelan DNA. By 1983, a

group at Indiana University was matching probe data to pedigrees, seeking a pattern. Luckily, the eighteenth of several hundred DNA probes, called G8, bound only the DNA of the sick people. G8 was linked to, and inherited with, the HD gene. Until the gene itself was discovered in 1993, this marker was the basis of presymptomatic testing.

The next step was to localize G8 to a chromosome. The researchers used hybrid rodent/human cells that contain only one human chromosome each. The hybrid cell that included G8 had human chromosome 4 and therefore, the HD mutation was in that chromosome. To extend the DNA sequence from the marker to the HD gene, the researchers located another probe that overlapped the first, then another probe that overlapped the second probe, and so on. This technique was called “chromosome walking.” Finally, a computer aligned the probes according to their sequence overlaps, creating a map of the extended DNA sequence. Where was the HD gene?

The area surrounding a probe on a chromosome can harbor many genes. Researchers first identified protein-encoding regions by looking for stretches of CGCGCG called “CpG islands,” which precede genes. That analysis showed that the half-million-base-long map could include 100 genes! Which were “candidate genes” for HD? The next clue was the phenotype. The researcher who had discovered that an expanding triplet repeat causes myotonic dystrophy (see figure 12.10) suggested that perhaps such a mutation lay behind HD, too, since both disorders affect movement.

The researchers indeed found a triplet repeat gene in the region that is longer in people who have HD and it is expressed in the brain. The journey from marker to mutation had taken a decade.

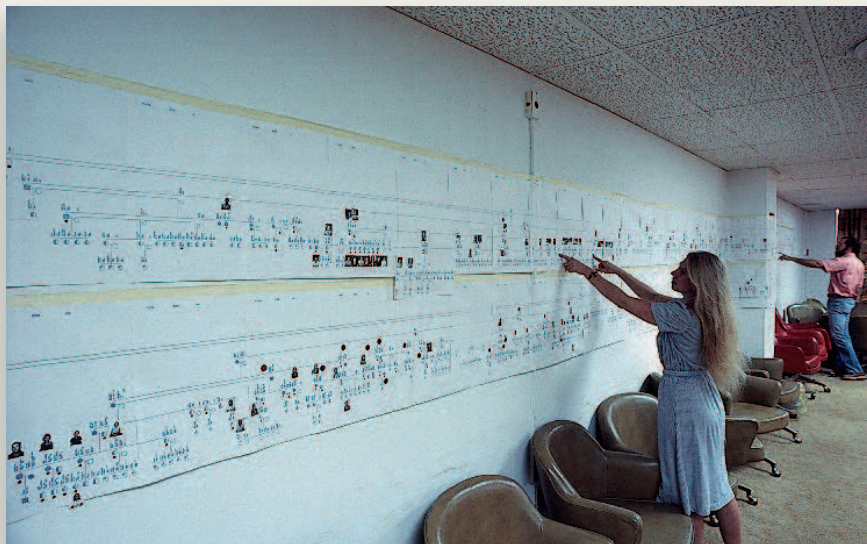


Figure 1 Nancy Wexler consults the enormous Venezuelan HD pedigree.

The Human Genome Project

The human genome project officially got underway in 1990, produced a “rough draft” in 2001, and a finished sequence in 2003 (see the Technology Timeline). Deciphering the functions of DNA sequences continues today, and has been complicated by the discovery that although much of the genome does not encode protein, much of it is nonetheless transcribed into RNA.

The idea to sequence the human genome occurred to many researchers at about the same time, but with different goals. It was first brought up at a meeting held by the Department of Energy (DOE) in 1984 to discuss the long-term population genetic effects of exposure to low-level radiation. In 1985, researchers meeting at the University of California, Santa Cruz,

called for an institute to sequence the human genome, because sequencing of viral genomes had shown that it could be done. The next year, virologist Renato Dulbecco proposed that the key to understanding the origin of cancer lay in knowing the human genome sequence. Later that year, scientists packed a room at the Cold Spring Harbor Laboratory on New York’s Long Island to discuss the feasibility of a human genome project. At first those against the project outnumbered those for it 5 to 1. The major fear was the shifting of goals of life science research from inquiry-based experimentation to amassing huge amounts of data.

A furious debate ensued. Detractors claimed that the project would be more gruntwork than a creative intellectual endeavor, comparing it to conquering Mt. Everest just because it is there. Practical benefits would be very far in the future. Some

Technology Timeline

EVOLUTION OF GENOME PROJECTS

1985–1988	Idea to sequence human genome suggested at several scientific meetings.
1988	Congress authorizes the Department of Energy and the National Institutes of Health to fund the human genome project.
1989	Researchers at Stanford and Duke Universities invent DNA microarrays.
1990	Human genome project officially begins.
1991	Expressed sequence tag (EST) technology identifies protein-encoding sequences.
1992	First DNA microarrays available.
1993	Need to automate DNA sequencing recognized.
1994	U.S. and French researchers publish preliminary map of 6,000 genetic markers, one every 1 million bases along the chromosomes.
1995	Emphasis shifts from gene mapping to sequencing. First genome sequenced: <i>Haemophilus influenzae</i> .
1996	Resolution to make all data public and updated daily at GenBank website. First eukaryote genome sequenced—yeast.
1998	Public Consortium releases preliminary map of pieces covering 98 percent of human genome. Millions of sequences in GenBank. Directions for DNA microarrays posted on Internet. First multicellular organism’s genome sequenced: roundworm.
1999	Rate of filing of new sequences in GenBank triples. Public Consortium and two private companies race to complete sequencing. First human chromosome sequenced (22).
2000	Microarray technology flourishes. First plant genome sequenced. Completion of first draft human genome sequence announced at White House.
2001	Two versions of draft human genome sequence published.
2003	Finished version of human genome sequence announced to coincide with fiftieth anniversary of discovery of DNA structure. Entire protein-encoding part of human genome available on DNA microarrays.
2005	Annotation of human genome sequence continues gradually. Number of species with sequenced genomes soars.
2007	Detailed analysis of a representative one percent of the human genome confirms that many genes have introns and reveals that most of the genome is transcribed. First individual human genome sequenced.
2008	First genome synthesized (<i>Mycoplasma genitalium</i>).
2010	Several human genomes sequenced.

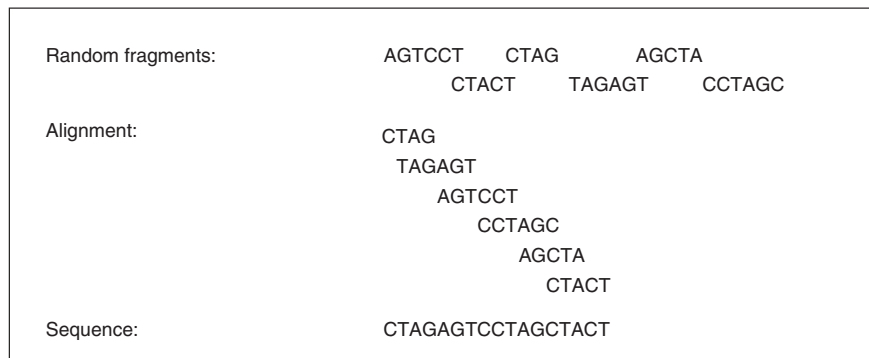


Figure 22.2 Deriving a DNA sequence. Automated DNA sequencers first determine the sequences of short pieces of DNA, or sometimes of just the ends of short pieces. Then algorithms search for overlaps. By overlapping the pieces, the software derives the overall DNA sequence.

researchers feared that such an unprecedented “big science” project would divert government funds from basic research and AIDS. Finally, the National Academy of Sciences convened a committee representing both sides to debate the feasibility, risks, and benefits of the project. The naysayers were swayed to the other side. In 1988, Congress authorized the National Institutes of Health (NIH) and the DOE to fund the \$3 billion, 15-year human genome project, which began in 1990 with James Watson at the helm. The project set aside 3 percent of its budget for the ethical, legal, and social issues program (ELSI). It has helped ensure that genetic information is not misused to discriminate.

Eventually, an international consortium and a private company, Celera Genomics, sequenced the human genome.

A series of technological improvements sped the genome project. In 1991, a shortcut called expressed sequence tag (EST) technology enabled researchers to quickly pick out genes most likely to be implicated in disease. This was a foreshadowing for future efforts to focus on the **exome**, which is the part of the genome that encodes protein. ESTs are cDNAs (see figure 19.5) made from the mRNAs in a cell type that is abnormal in a particular illness. Also in that year, DNA microarrays were developed to display short DNA molecules, which became important in sequencing (tiling arrays) as well as in assessing gene expression (expression arrays).

Development of computer algorithms eased the assembly of many short pieces of DNA with overlapping end sequences into longer sequences (**figure 22.2**). (The next section discusses the actual DNA sequencing.) When the project began, researchers cut several genomes-worth of DNA into overlapping pieces of about 40,000 bases (40 kilobases), then randomly cut the pieces into small fragments. The greater the number of overlaps, the more complete the final assembled sequence. The sites of overlap had to be unique sequences, found in only one place in the genome. Overlaps of repeated sequences found in several places in the genome could lead to more than one derived overall sequence—a little like searching a document for the word “that” versus searching for an unusual word, such as “dandelion.” Searching for “dandelion” is more likely to lead to a specific part of a document, whereas “that” may occur in several places—just like repeats in a genome. The use of unique sequences is why the human genome project did

not uncover copy number variants. For example, the sequence CTACTACTA would only appear as CTA. Researchers did not at first appreciate the fact that repeats are a different form of information than DNA base sequences.

Two general approaches were used to build long DNA sequences (**figure 22.3**). The “clone-by-clone” technique used by the U.S. government-funded group aligned pieces one chromosome at a time. The “whole genome shotgun” approach used at Celera Genomics shattered the entire genome, then used a computer algorithm to identify overlaps and align them to derive a continuous sequence. The task can be compared to cutting the binding off a large book, throwing it into the air, and reassembling the dispersed pages in order. A “clone-by-clone” dismantling of the book would divide it into bound chapters. The whole genome shotgun approach would free every page. Whole genome shotgunning is faster, but it misses some sections (particularly repeats) that the clone-by-clone method detects.

Technical advances continued. In 1995, DNA sequencing became automated, and software was developed that could rapidly locate the unique sequence overlaps among many small pieces of DNA and assemble them into a continuous sequence, eliminating the need to gather large guidepost pieces.

In 1999, the race to sequence the human genome became intensely competitive. The battling factions finally called a truce. On June 26, 2000, Craig Venter from Celera Genomics and Francis Collins, representing the International Consortium (and now director of the NIH), flanked President Clinton in the White House rose garden to unveil the “first draft” of the human genome sequence. The milestone capped a decade-long project involving thousands of researchers, which was the culmination of a century of discovery. The historic June 26 date came about because it was the only opening in the White House calendar! In other words, the work was monumental; its announcement, staged.

Figure 22.4 is an overview of genome sequencing. **Table 22.1** lists websites that have genome sequence information.

Looking back, was sequencing the human genome valuable, or was it indeed like climbing Mt. Everest just because it is there? Availability of the sequence has greatly accelerated

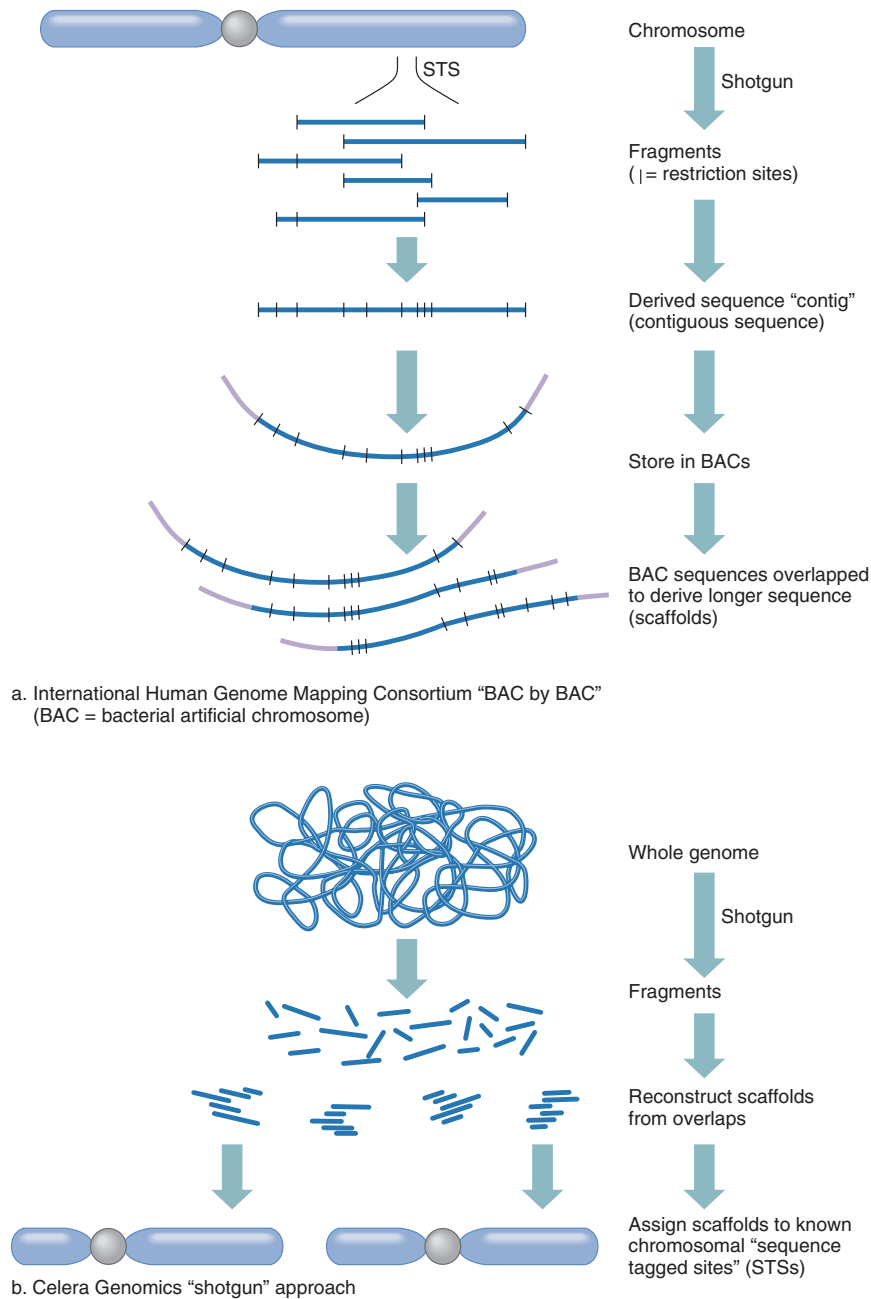


Figure 22.3 Two routes to sequencing the human genome. (a) The International Consortium began with known chromosomal sites and overlapped large pieces, called contigs, that were reconstructed from many small, overlapping pieces. "STS" stands for "sequence tagged site," which refers to specific known parts of chromosomes. A BAC is a cloning vector that uses bacterial DNA. (b) Celera Genomics shotgunned several copies of a genome into small pieces, overlapped them to form scaffolds, and then assigned scaffolds to known chromosomal sites. They used some Consortium data.

gene discovery beyond the slow disease-by-disease approach of the pre-genome era, but gene expression profiling (chapter 11) and genome-wide association studies (chapter 7 and others) have also impacted health care and revealed human diversity. In addition, the early focus on stating "the" number of human genes seems misplaced in hindsight. Just as learning the words

of a language is not the same as reading stories in that language, it is the interactions among genes that are important, not their number. But if we have learned anything, it is that we can't predict future utility of discoveries. **Reading 22.2** describes some of the information found in sequencing the first three genomes of specific individuals.

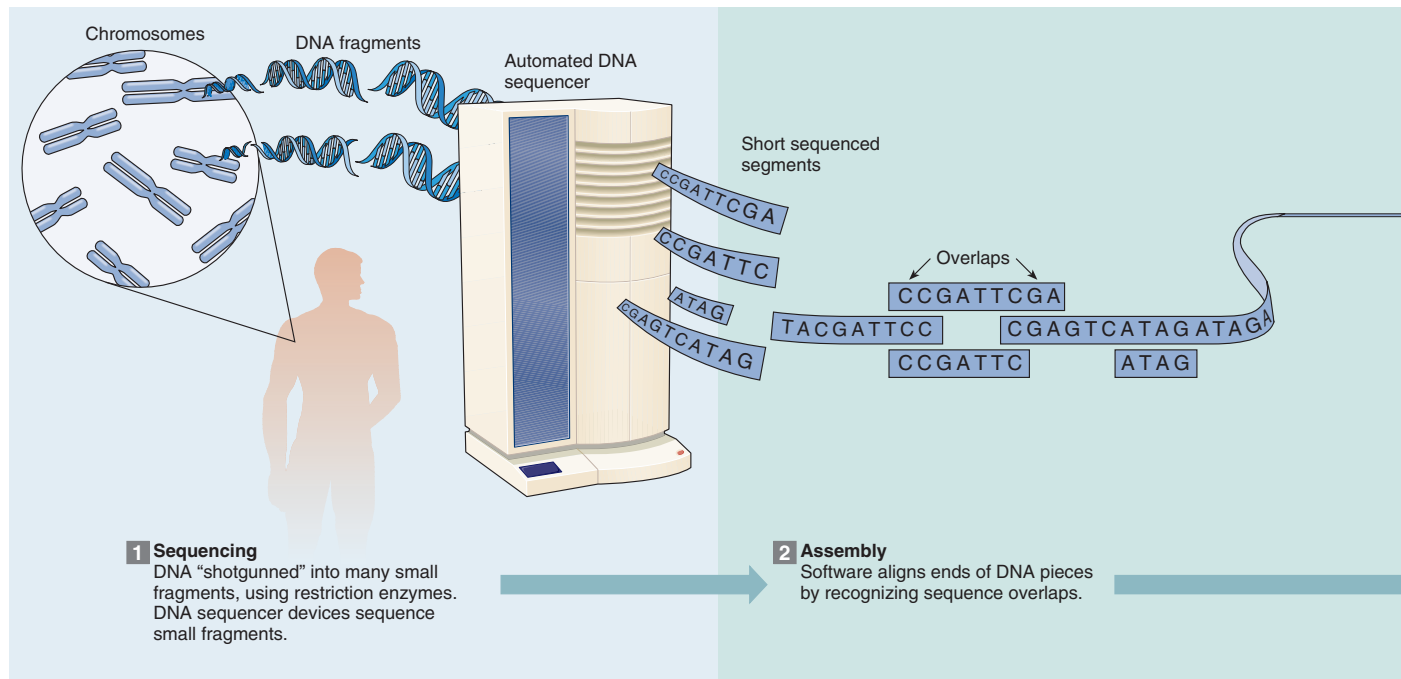


Figure 22.4 Sequencing genomes. The "shotgun" approach to genome sequencing overlapped DNA pieces cut from several copies of a genome, then assembled the overall sequence. Newer techniques use microfluidics and nanomaterials to sequence DNA.

Table 22.1 Public Genome Databases

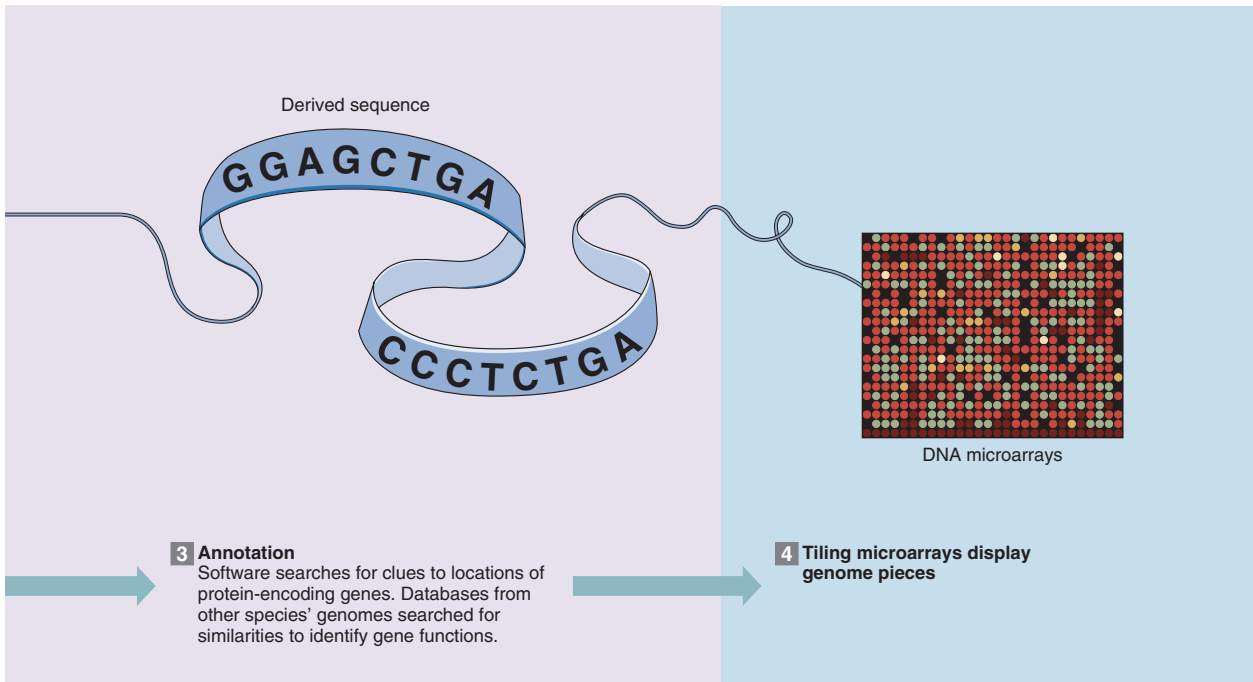
Organization	Website
GenBank	http://www.ncbi.nlm.nih.gov/Genbank/
European Molecular Biology Laboratory (EMBL)	www.ebi.ac.uk/embl/index.html
DNA Data Bank of Japan	www.ddbj.nig.ac.jp
University of California, Santa Cruz Genome Browser	www.genome.ucsc.edu/
National Center for Biotechnology Information Map Viewer	www.ncbi.nlm.nih.gov/mapview/
National Human Genome Research Institute, NIH	www.genome.gov/
SNP Database	www.ncbi.nlm.nih.gov/projects/SNP
U.S. Department of Energy Genomes to Life	http://genomicsgsl.energy.gov/
Genomes OnLine Database (GOLD)	http://www.genomesonline.org/

Key Concepts

- Human genome sequencing was built on linkage and cytogenetic information from decades of work.
- Positional cloning located specific disease-causing genes in families.
- The idea to sequence the human genome emerged in the mid-1980s with several goals. The project officially began in 1990.
- Genome sequencing cuts several copies of a genome, sequences the pieces, then uses algorithms to overlap the pieces.
- Clone-by-clone sequencing assembled chromosomes individually. Whole genome shotgunning shattered the entire genome and rebuilt it.

22.2 DNA Sequencing and Genome Synthesis

Several new ways to sequence DNA are available today. **Table 22.2** indicates the great improvement in DNA sequencing efficiency with time.



3 Annotation
Software searches for clues to locations of protein-encoding genes. Databases from other species' genomes searched for similarities to identify gene functions.

4 Tiling microarrays display genome pieces

One of the original methods for sequencing DNA, invented in 1977 by Frederick Sanger, is still widely used and was quite brilliant in concept. The Sanger method generates a series of DNA fragments of identical sequence that are complementary to the DNA sequence of interest. These fragments differ in length from each other by one end base, as follows:

Sequence of interest: T A C G C A G T A C
Complementary sequence: A T G C G T C A T G
Series of fragments:



Note that the entire complementary sequence appears in the sequence of end bases of each fragment. The complement of the gene of interest is cut into a collection of pieces, differing in the end bases, which are distinguished with a radioactive or fluorescent label. That is, A, T, C, and G are labeled with different fluorescent colors. Then the fragments are separated by size. Once the areas of overlap are aligned, reading the labeled end bases of the pieces in size order reveals the sequence of the complement, from which the sequence of interest is derived. **Figure 22.5** shows how DNA sequence data derived from the Sanger method appear in scientific papers, and **figure 22.6** shows how the sequence is read from the end bases.

Newer approaches to DNA sequencing use a microfluidics environment, which is a small, fluid-filled chamber. One

Year	# DNA Bases Sequenced/Day
1986	1,000
1995	15,000
1997	500,000
1998	1 million
2005	80 million
2009	8.8 billion

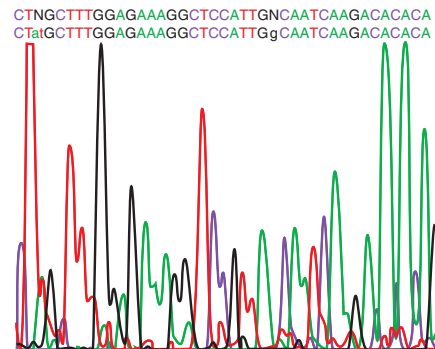


Figure 22.5 DNA sequence data. In automated DNA sequencing, a readout of sequenced DNA is a series of wavelengths that represent the terminal DNA base labeled with a fluorescent molecule.

Reading 22.2

The First Three Human Genome Sequences

"The diploid genome sequence of an individual human," *PLoS Biology*, October 2007 (Craig Venter)

"The complete genome of an individual by massively parallel DNA sequencing," *Nature*, April 17, 2008 (James Watson)

"The Diploid Genome Sequence of an Asian Individual," *Nature*, November 6, 2008 ("YH")

The genomes sequenced in the human genome projects of 2001 were composites of different individuals. The first two genomes of specific individuals to be sequenced—of genome research pioneers Craig Venter and James Watson—yielded few medical surprises. Instead, they showed that we had greatly underestimated genetic variation by focusing only on the DNA sequence. The numbers of copies of short sequences—copy number variants, or CNVs—contribute significantly to genetic variation.

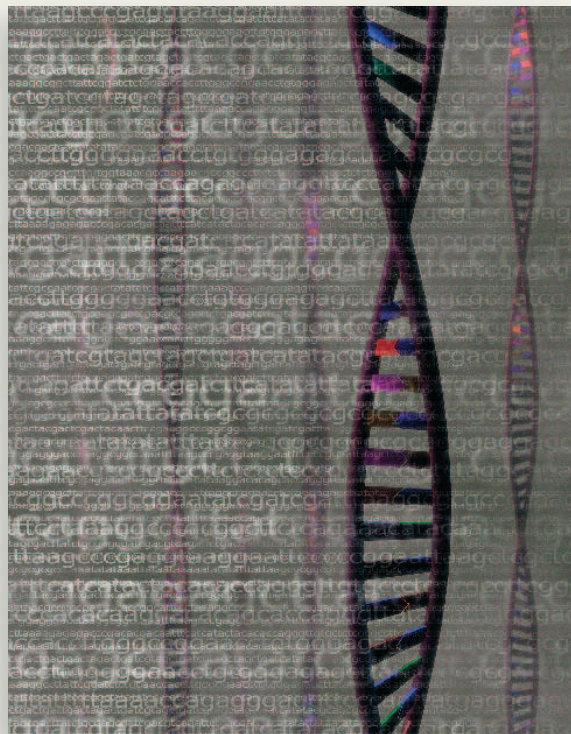
"Back in 2001, we thought we differed from chimps by 1.27% of our genomes. Now we know that we differ from each other by as much as 1 to 3%. If we count all the differences, we are about 5 to 6% from the chimp. In the way we put sequences in public databases, we lost the insertions and deletions," said Venter to an American Society of Human Genetics meeting after he'd had a year to think about what his personal genome sequencing had revealed. It wasn't much that he didn't already know from his family history and personal experience.

Venter has gene variants associated with increased risk of Alzheimer disease and cardiovascular disease. He has alleles for dry earwax, blue eyes, lactose intolerance, a preference for activities in the evening, and a tendency toward antisocial behavior, novelty seeking, and substance abuse. Not to his great surprise, he is a fast metabolizer of caffeine. "I can have two double lattes and wash it down with a Red Bull and not be affected by it," he said.

James Watson, according to his genome sequence, is a heterozygote for a dozen rare recessive disorders, including a glycogen storage disease, two eye conditions, and a DNA repair disorder, and he is at elevated risk for twenty other disorders. Science journals deemed Watson's results "of thin clinical value" and yielding "few biological insights." Said Richard Gibbs, one of the researchers who sequenced Watson's genome, "We tried genetic counseling on Jim, but it was a failure." Yet Watson and Venter differ in inherited drug responses, supporting the value of pharmacogenetics/genomics (discussed in chapter 20). Said Venter, "You probably wouldn't suspect this based on our appearance—we are both bald, white scientists."

The third person to have his genome sequenced was called, simply, "YH." He is Han Chinese, an East Asian population that accounts for 30 percent of modern humanity. He has no inherited diseases in his family, but his genome includes 116 gene variants that cause recessive disorders, as well as many risk alleles. He shares with Craig Venter a tendency to tobacco addiction and high-risk alleles for Alzheimer disease.

An overall comparison of the first three genome sequences of individuals provides a peek at our variation. Each man has about 1.2 million SNPs, but a unique collection. Each has only .20 to .23 percent of SNPs that are nonsynonymous, meaning that they alter an encoded amino acid, and the men share only 37 percent of these more meaningful SNPs. The math indicates, therefore, that about .07 percent of our SNPs may affect our phenotypes.



What will we learn from our personal genome sequences?

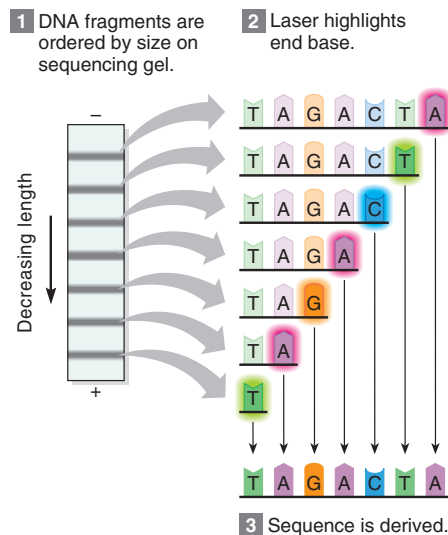


Figure 22.6 Reading a DNA sequence. A computer algorithm detects and records the end base from a series of size-ordered DNA fragments.

technique sequences many 100-base-long single-stranded DNA pieces that are attached to tiny beads in a water-oil mixture. A laser reads off fluorescently tagged bases that are added, according to the complementary base pair rules, as nucleotides stream past the strands. The method, called 454 sequencing, was invented by a father seeking a way to quickly sequence his newborn son's genome. The technique can sequence 20 million bases in about four and a half hours. Even newer approaches use nanomaterials to distinguish how each of the four nucleotide bases disrupts an electrical field as a DNA molecule passes through nanopores at about 1,000 bases per second. Various "nano" approaches have been developed to rapidly sequence DNA.

If it is possible to *sequence* a genome, it should also be possible to *synthesize* a genome. Researchers at the J. Craig Venter Institute did this in 2008 for the 582,970 bases of *Mycoplasma genitalium*, a tiny bacterium. The researchers worked with 5,000- to 7,000-base-long "cassettes" of the known sequence, synthesizing in the laboratory sets of pieces 100 bases long that overlapped at the ends. They then delivered the pieces into other cells (*E. coli* or yeast), using vectors called bacterial artificial chromosomes (BACs). Natural recombination joined the pieces, building first a quarter of the genome, then half, then three-quarters, and finally the entire genome.

The goal of constructing the bacterial genome was not to create life, and so the researchers did not use the assembled genetic instructions to make a functional microorganism. However, the ability to create a genome is a very new research approach. Geneticists have traditionally identified a gene's function by mutating it in a model organism and observing what goes wrong. Synthesizing a genome is instead a way to identify a gene's function by creating it from chemicals.

Key Concepts

1. In the Sanger method of DNA sequencing, complementary copies of an unknown DNA sequence are cut into different-sized pieces differing from each other by an end base. The pieces are overlapped by size and the labeled end bases read off.
2. Newer techniques use short DNA sequences and nanomaterials.
3. Researchers have built the first genome.

22.3 Ways of Looking at Genomes

If Wikipedia were transcribed into old-fashioned printed volumes, the information would fill many books. So it is with a human genome. Just as you wouldn't read all of Wikipedia to learn about toenail fungus or opera, not all of the 3.2 billion bases of a human genome hold information that is useful to a particular person in a particular circumstance.

Long before the human genome was sequenced, researchers' thoughts turned to what they would do with the information. Today, genome information is being accessed and analyzed at several levels:

- Between and among species, for views on evolution.
- In a representative 1 percent to discover how a genome works and the types of information held in DNA sequence and organization.
- In subsets of sequence, such as the protein-encoding portion.

Comparative Genomics

Hundreds of species have had their genomes sequenced. The first were viruses and bacteria, because genome shotgunning works best on small genomes. Then the genomes of our closest relatives were sequenced—mice, rats, chimps and our favorites, cats and dogs. Most informative, however, have been the genomes of species that represent evolutionary crossroads. These are organisms that introduced a new trait or were the last to have an old one.

In comparing genomes of modern species, researchers infer evolutionary relationships from DNA sequences that are conserved and presumably selected through time. **Figure 22.7** shows one way of displaying short sequence similarities, called a pictogram. DNA sequences from different species are aligned, and the bases at different points indicated. A large letter A, C, T or G indicates, for example, that all species examined have the same base at that site. A polymorphic site, in contrast, has different bases for different species.

The field of comparative genomics uses conserved sequences to identify biologically important genome regions,

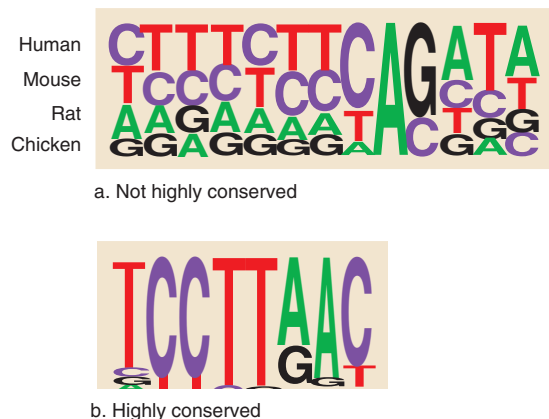


Figure 22.7 A pictogram indicates conservation of sequence. Genomes can be compared site by site. These pictograms are for short sequences in corresponding regions of the human, mouse, rat, and chicken genomes. A large letter means that all four species have the same base at that site. If four letters appear in one column, then the species differ. Pictogram (a) is not highly conserved; (b) is.

assuming that persistence means evolutionary success. But there are exceptions—conserved sequences with no apparent function in humans. Either we haven’t discovered the functions, or genomes include “raw material” for future functions. **Figure 22.8** presents a few organisms whose genomes have been sequenced. Following are examples of the types of information inferred from conserved DNA sequences.

The minimum gene set required for life The smallest microorganism known to be able to reproduce is *Mycoplasma genitalium*. It infects cabbage, citrus fruit, corn, broccoli, honeybees, and spiders, and causes respiratory illness in chickens, pigs, cows, and humans. Researchers call its tiny genome the “near-minimal set of genes for independent life.” Of 480 protein-encoding genes, 265 to 350 are essential. Considering how *Mycoplasma* uses its genes reveals the fundamental challenges of being alive. This was the first organism to have its genome synthesized.

Fundamental distinctions among the three domains of life *Methanococcus jannaschii* is a microorganism that lives at the bottom of 2,600-meter-tall “white smoker” chimneys in the Pacific Ocean, at high temperature and pressure and without oxygen. As archaea, these cells lack nuclei, yet replicate DNA and synthesize proteins in ways similar to multicellular organisms. The genome sequence confirmed that this organism represents a third form of life.

The simplest organism with a nucleus The yeast *Saccharomyces cerevisiae* is single-celled with only about 6,000 genes, but a third of them have counterparts among mammals, including at least seventy genes implicated in human diseases.

Understanding what a gene does in yeast can provide clues to how it affects human health, such as counterparts of mutations in cell cycle control genes in yeast causing cancer in humans.

The basic blueprints of an animal The genome of the tiny, transparent, 959-celled nematode worm *Caenorhabditis elegans* is packed with information on what it takes to be an animal. Thanks to researchers who, in the 1960s, meticulously tracked the movements of each cell as the animal developed, much of the biology of this organism was already known before its 97 million DNA bases were revealed late in 1998. The worm’s signal transduction pathways, cytoskeleton, immune system, apoptotic pathways, and brain proteins are very similar to our own. Curiously, the fruit fly (*Drosophila melanogaster*) genome has 13,601 genes, fewer than the 18,425 in the much simpler worm. Of 289 disease-causing genes in humans, 177 have counterparts in *Drosophila*. The fly is a model for testing new treatments.

Life on land Before 450 million years ago, according to fossil evidence, life was confined to the seas, where it was abundant and diverse. Algae, microorganisms, and jawless fishes shared the depths. The first organisms to colonize land were the mosses, and for this reason, the genome of the modern moss *Physcomitrella paten* was sequenced. Mosses lack stems and leaves and have only a few types of differentiated cells. They dominated landscapes until plants that had seeds and vessels evolved some 200 million years later.

By the time animals ventured onto land, plants had already taken root. Sea residents whose descendants were probably among those first land dwellers were the lobe-finned fishes, which have fleshy, strong fins that could have evolved into limbs. Two types of lobe-finned fishes persist today and resemble their fossilized forms—the lungfishes and two species of coelacanths. Because the lungfish genome is huge, researchers analyzed the smaller coelacanth genome. Once thought to be extinct, coelacanths today live in the Indian Ocean. Information in the coelacanth genome may reveal the traits necessary for the evolution of the tetrapods—vertebrates with four limbs.

From birds to mammals The sequencing of the chicken genome (figure 22.8b) marked a number of milestones—the first agricultural animal, the first bird, and, as such, the first direct descendant of dinosaurs. The genome of the red jungle fowl *Gallus gallus* is remarkably like our own, minus many repeats, but its genome organization is intriguing. Like other birds, fishes, and reptiles, but not mammals, the chicken genome is distributed among very large macrochromosomes and tiny microchromosomes. Repeats may have been responsible for the larger sizes of mammalian chromosomes.

From chimps to humans Most comparisons of the human genome to those of other species seek similarities. But comparisons of our genome to the genome of the chimpanzee *Pan troglodytes* aim to find the opposite—genetic differences may refine our knowledge about what makes humans unique, discussed in Reading 16.1. Our genomes differ

<p>Honey Bee (<i>Apis mellifera</i>)</p> <p>~300 million bases</p>		<p>Red jungle fowl (<i>Gallus gallus</i>)</p> <p>~1 billion bases</p>	
<p>■ Agriculture: honey</p> <p>■ Animal societies</p> <p>■ Compare to other insect genomes</p> <p>■ Ecology: Compare to Africanized bees in southwestern U.S.</p> <p>a.</p>		<p>■ Evolution: Dinosaur descendant; conserved control sequences</p> <p>■ Good model organism. Can study early development in eggs, and aging</p> <p>■ Same number of genes as humans, but genome 1/3 the size</p> <p>■ Agriculture: Identify genes that limit need for drugs in feed</p> <p>■ Medicine: Carries avian flu virus</p> <p>b.</p>	
<p>Tammar wallaby (<i>Macropus eugenii</i>)</p> <p>~3.6 billion bases</p>		<p>Dog (<i>Canis familiaris</i>)</p> <p>~2.5 billion bases</p>	
<p>■ Evolution: Marsupials (pouched mammals split from placental mammals ~130 million years ago)</p> <p>■ Perpetually pregnant; give birth on same day each year</p> <p>■ 1 million on Kangaroo Island, Australia</p> <p>c.</p>		<p>■ Evolution: Extreme artificial selection created 300+ breeds</p> <p>■ 400+ diseases from founder effect and inbreeding</p> <p>■ Medicine: Diseases in humans too: rheumatoid arthritis, cancers, heart and eye disorders, deafness</p> <p>■ Biotechnology: Pioneered diabetes treatment and bone marrow transplant</p> <p>d.</p>	

Figure 22.8 A sampling of animal genomes.

by 1.2 percent, equaling about 40 million DNA base substitutions. Within those differences, as well as copy number distinctions, may lie the answers to compelling medical questions. Our cancer rates differ; humans are susceptible to malaria, but chimps are not; humans develop Alzheimer disease, and chimps do not; and the course of HIV is deadlier than the chimp version, SIV.

An international consortium is cataloging species according to selected short genome regions that vary among species but not much within species. Chapter 1 discusses this genetic “barcoding.”

A Representative One Percent

One way of examining the human genome is to randomly sample a small portion of it, and catalog what the DNA sequences do. The ENCODE (Encyclopedia of DNA Elements) project is

considering a representative 1 percent of the human genome. The coverage, spread among the chromosomes, amounts to 30 million of our 3 billion DNA bases. The DNA sequences are from known, well-studied genes as well as uncharted territory, with an initial goal of deriving a “parts list” of that 1 percent.

The ENCODE project is filling in some gaps between the single-gene protein-encoding approach that dominated genetics for decades and evolutionary clues from comparing the genomes of many species. So far, results have confirmed and extended a complexity that researchers had begun to suspect with the discovery of introns in the late 1970s (table 22.3). *The old view:* The human genome is a static collection of individual protein-encoding genes, acting alone, mired in a sea of “junk” sequences. *The new view:* Most, if not all, of the human genome consists of classes of DNA that have different types of functions that interact through space and time to control cellular

Table 22.3 ENCODE Results

Analysis of 1 percent of the human genome reveals that:

- Most of the genome is transcribed.
- DNA sequences previously thought not to be transcribed actually are, and some overlap exons.
- The genome has more promoters, and therefore more ways to build genes from its parts, than suspected.
- Control sequences are not only at the starts of genes—they are all over.
- Chromatin structure controls DNA replication as well as transcription.
- 5% of the human genome is highly conserved in all mammals.
- Functional parts of the genome may vary among species. These sequences may represent a DNA store that may serve as “a warehouse for evolution” that may be species-specific.

activities. DNA sequences encode amino acid sequences, but they also control each other in intricate networks of interactions. The 1960s-era view of one-gene/one-protein was clearly a great oversimplification of the way the human genome works.

Probing the Human Exome

The exome includes DNA sequences that encode the parts of genes that are transcribed into RNA and then translated into protein. This is the collection of exons in a genome—what geneticists have been studying since the middle of the twentieth century, as they matched phenotypes to first chromosome abnormalities and then rare mutations in single genes. The human exome includes about 180,000 exons. Exploring the exome gene-by-gene, as in the past, is a bottoms-up approach; having an entire human genome sequence at hand provides a top-down approach. This is what researchers did with the first sequenced human genome, termed “HuRef” for human reference genome. The analysis of the HuRef exome used many databases and reports and included as variants SNPs and indels, which are *insertions* or *deletions* of bases. Indels can disrupt the reading frame if they are not a multiple of three, or can shorten a protein by introducing a stop codon.

Recall from chapter 10 that nonsynonymous SNPs are single nucleotide polymorphisms—sites of single base changes—that alter the sequence such that a different amino acid is specified. For example, a change of CCG to CCC does not change the amino acid specified, which is proline (see figure 10.12). This is a synonymous change. However, changing CCG to CAG alters the specified amino acid to glutamine. Whether or not a nonsynonymous SNP affects the phenotype depends on whether or not it is in a part of the protein that affects its function.

Analysis of the HuRef exome confirms and extends the observation that very little of our genome encodes protein. Of approximately 12,500 variants in the exome, 10,400 are nonsynonymous SNPs, of which about 20 percent are rare and most recessive. The remaining 80 percent are in genes associated with

common traits or disorders. Of the 700 indels, about half are multiples of three, which might not disrupt the reading frame sufficiently to affect the functioning of the encoded protein.

All in all, of the 12,500 or so variants in protein-encoding genes in this sample exome, only about 1,500 alter protein function, thereby potentially affecting the phenotype. This is apparently the pool from which a person’s genetic load—collection of dangerous alleles—comes. Perhaps so few genetic diseases have been identified because there really aren’t as many as early estimates of human gene number suggested. **Table 22.4** presents an overview of exomic diversity between the first two human genomes sequenced.

Many researchers study subsets of exomes—that is, classes of genes whose encoded proteins are functionally related. The lipidome, for example, includes genes whose protein products participate in the synthesis or breakdown of lipids (fats). The secretome includes proteins that are secreted from the cell. “Omics” already mentioned include the diseaseome, metagenome, and microbiome (chapter 1), the transcriptome (chapter 11), and the pharmacogenome (chapter 20). “Omics” are becoming more and more specific. For example, the “kinome” includes all of the genes that encode a type of enzyme called a kinase.

Even as “omics” proliferate and fuel gene discovery, genetic research is coming full circle, as investigators continue to compile catalogs of the mutations that cause specific disorders in “locus-specific databases.” A global effort called the Human Variome Project is bringing together the hundreds of existing databases so that researchers can be aware of each other’s work. Many of the databases represent the efforts of disease-specific organizations, which are often started and run by parents of children who have inherited disorders.

It will be interesting to see how the genome-level studies like the HuRef exome and the locus-specific databases come together. The genome-wide association studies will help to bridge the gap. These studies highlight genome regions where sequence variants track with interesting traits or conditions, leading to identification of causative genes. The puzzle that is the human genome is gradually coming together.

Table 22.4 The Exomes of J. Craig Venter and James Watson

Genome Feature	Venter	Watson
# nonsynonymous SNPs	10,389	10,569
# novel nonsynonymous SNPs	772 (7%)	1,573 (15%)
% that likely affect protein function	14%	20%
# coding indels of at least 2 bases	423	345

A nonsynonymous SNP is a single base change that alters the encoded amino acid.

Novel SNPs are those not in the SNP database (<http://www.ncbi.nlm.nih.gov/projects/SNP/>).

Indels are insertions and deletions.

Data from P. C. Ng et al., “Genetic variation in an individual human exome,” *PLoS Genet.* 2008 August; 4(8):e1000160, Table 4)

Key Concepts

1. Comparative genomics infers evolutionary relationships from conserved DNA sequences.
2. The ENCODE project is taking an in-depth look at 1 percent of the human genome.
3. The exome is a collection of the exons in the human genome. Only a small proportion of its variation affects the phenotype. “Omics” are subsets of exomes.
4. Locus-specific databases compile mutations in single genes. Researchers study both entire genomes and single genes.

22.4 Personal Genome Sequencing

With the turn of the century, genetics was catapulted from a somewhat obscure science to an increasingly familiar field with many practical applications, while providing a new view of humanity in the larger context of all life on Earth. *Bioethics: Choices for the Future* on page 444 presents some viewpoints on how genome information should be used.

On a personal level, human genome information will focus on the healthy as well as the ill, to better understand disease. Large-scale investigations of the genomes of remarkably healthy centenarians—those past age 100—will reveal the spectrum of gene variants that keep them well (see Reading 3.1). Such projects are not attempts to define a “perfect” genetic blueprint, but aim to develop guidelines that might suggest new types of treatments based on how protective gene variants function and interact.

Identification of disease-causing genes has proceeded for half a century and will continue, but with unprecedented precision. At the same time, each genetically influenced disease in an individual will be considered in the context of the entire genome. Future treatments based on genome information will not be limited to correcting or circumventing genetic flaws, or supplying missing proteins. Discovering which gene variants contribute to which diseases, and how they do so, will enable us to identify risk factors that are easier to control—those from the environment.

You might learn your own genetic information in several ways: from a test for a single-gene mutation, from a selected list of associations chosen from a direct-to-consumer genetic testing website, or from sequencing of your exome or entire genome. To help decide whether to have your genome sequenced, imagine how the information might be used, or misused:

- Despite existing legislation, health insurance might come to be based on which diseases and risks you have inherited.
- A diagnosis of cancer, prediction of its course, or determining which treatments will work best, might be done in seconds following a genome scan.
- Gene variants that affect intelligence might be used in college admission decisions or in class placements.
- Gene variants that affect strength and endurance might be considered in tryouts for athletic teams or used to determine whether some competitors have unfair genomic advantages.
- People might consider genome information in choosing partners.

The ability to easily sequence our genomes continues a long-held curiosity about our genetic selves. Since the dawn of humanity, people have probably noted inherited traits, from height and body build, to hair and eye color, to talents, to behavioral quirks, to illnesses. Genetics provides the variety that makes life interesting.

The science of genetics grew out of questions surrounding plant and animal breeding. It became human-oriented in the mid-twentieth century with the recognition that certain characteristics and conditions run in families, sometimes recurring with predictable frequencies. Today, genetics and genomics impact many areas.

In the near future, you might

- be offered a DNA microarray test to diagnose or treat a medical condition.
- serve on a jury and be asked to evaluate DNA profiling evidence.
- take a panel of genetic tests before trying to have a child.
- seek preimplantation genetic diagnosis to ensure that your child does not inherit a particular genotype.
- help a parent or other loved one through chemotherapy, with some assurance, thanks to DNA testing, that the most effective drugs will be tried first.
- eat a genetically modified fruit or vegetable (you likely have already done this).
- receive a body part from a pig, with cell surfaces matched to your own.

The list of applications of genetic and genomic technologies is long and ever-expanding. I hope that this book has offered you glimpses of the future and prepared you to deal personally with the choices that genetic technology will present to you. Let me know your thoughts!

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Bioethics: Choices for the Future

The \$1,000 Genome

Nature Genetics magazine asked readers to comment on how personal genome sequencing should be used once the cost falls. The many answers fell into several categories:

- **Health.** Sequence a family's genomes and use the information to guide lifestyle choices, screening tests for diseases, and taking drugs to stall or lessen severity of specific symptoms.
- **History.** Sequence genomes of people throughout the world who live in isolated, rural villages to trace ancient migration pathways.
- **Mutation.** Sequence the genomes of many cells from a single individual to reveal somatic mutations; from sperm or oocytes to shed light on meiosis. Sequence the genomes of monozygous twins and catalog health distinctions to isolate environmental influences. Compare genomes of parent-child pairs to reveal new mutations.
- **Cancer.** Sequence the genomes of cancer cells as the disease progresses in an individual to reveal genomic changes.
- **Benefit of sequencing.** Follow two groups, one of which has its members' genomes sequenced, and one not, to see if the knowledge translates into actions that improve health or extend life.
- **Less-tangible traits.** Sequence genomes to examine the genetic bases of genius, beauty, happiness, memory, personality, and behavior.
- **Evolution.** Sequence others—bacteria, crops, chimps, cats, dogs, or everyone and everything.

Perhaps the most valuable suggestion, summarizing many peoples' desire to dissect human diversity, came from David Goldstein at

Duke University: use the \$1,000 genome to learn “the genetics of normality.”

Questions for Discussion

1. Under what circumstances should a parent allow a young child's genome to be sequenced?
2. How can we best protect private genomic information?
3. Compare and contrast the accessibility of personal genome sequencing to that of the Internet or a cell phone. How would service contracts work to continually update relevant genetic discoveries?
4. What lessons learned from the Human Genome Project can be used to guide ethical use of personal genome information?
5. If there is tremendous demand for personal genome sequencing, how should requests be prioritized?
6. In medical practice, an incidentaloma is a test result that a physician wasn't looking for, like finding a cancerous tumor on an X ray of a broken bone. Discuss how personal genome sequencing could provide “too much information” in the form of incidentalomas.
7. A human exome can be sequenced for about \$10. Would sequencing an entire genome justify the difference in cost from doing just the exome, in terms of the value of the provided information?
8. Should genetic researchers accept payment as “sequence consultants” to help citizens understand the information in their genomes?
9. What would you do with knowledge of your own genome sequence?

Summary

22.1 From Genetics To Genomics

1. Genetic maps have increased in detail and resolution, from cytogenetic and linkage maps to physical and sequence maps.
2. Positional cloning discovered individual genes by beginning with a phenotype and gradually identifying a causative gene, localizing it to part of a chromosome.
3. The human genome project began in 1990 under the direction of the DOE and NIH. Technological advances sped the sequencing.
4. Several copies of a genome are cut and the pieces sequenced, overlapped, and aligned to derive the continuous sequence. For the human genome, the International Consortium used a chromosome-by-chromosome approach and Celera Genomics used whole genome shotgunning.
5. Since the sequencing of the human genome, other types of genome information are recognized as being equally or more important.

22.2 DNA Sequencing and Genome Synthesis

6. In the Sanger method of DNA sequencing, DNA fragments differing in size and with one labeled end base are aligned, and the sequence read off from the end bases.
7. Newer sequencing methods use microfluidics and nanomaterials.
8. Researchers synthesized the genome of a small bacterium.

22.3 Ways of Looking at Genomes

9. Identifying conserved regions among genomes of different species reveals some genes with vital functions.
10. Analysis of a representative one percent of the human genome showed that nearly all DNA is transcribed and that gene function is more complex than was thought.
11. The **exome** is the collection of exons in the human genome. Very little of the variation in the exome affects the phenotype. Other “omes” consider subsets of the exome.

22.4 Personal Genome Sequencing

12. Comparing the genomes of healthy people to those of people with certain diseases will reveal how gene combinations affect health, permitting recognition of controllable environmental factors.

13. Sequencing personal genomes is underway. The information can be used in various ways.

www.mhhe.com/lewisgenetics9

Answers to all end-of-chapter questions can be found at www.mhhe.com/lewisgenetics9. You will also find additional practice quizzes, animations, videos, and vocabulary flashcards to help you master the material in this chapter.



Review Questions

- Distinguish between a gene and a genome.
- How did family linkage patterns and chromosomal aberrations lay the groundwork for sequencing the human genome?
- Describe how the four levels of genetic maps differ.
- Explain how positional cloning was used to map the gene for Huntington disease.
- Why was there initial disagreement over whether the human genome should be sequenced?
- Why must several copies of a genome be cut to sequence it?
- Explain how individuals, such as James Watson and Craig Venter, can vary by 1 to 3 percent in their genome sequence, but not nearly that much in the amount of variation that affects the phenotype.
- Genomes of several bacteria have been synthesized in the lab, from what Craig Venter jokingly calls "four bottles of chemicals." What would be the next step in actually creating life?
- Explain how conserved DNA sequences hold information about evolution.
- List four types of information revealed in comparing genomes of different species.
- What has the ENCODE project revealed about the human genome?
- Why do researchers think that it might be more important to sequence the exome than an entire human genome?
- Explain why nonsynonymous SNPs can affect a phenotype but synonymous SNPs do not.
- If you have your genome sequenced, what will you do with the information?
- Why were Craig Venter and James Watson not terribly upset at the disease-related gene variants revealed in their genomes?

Applied Questions

- When the idea of sequencing the human genome was first discussed, some researchers thought it would be too straightforward and boring, and that amassing large amounts of information was not creative. Discuss one way that genome sequencing has turned out to be more complicated and/or more creative than anticipated.
- Suggest a species you believe should have its genome sequenced, and what information you think the sequence might reveal.
- Restriction enzymes break a sequence of DNA bases into the following pieces:
TTAATATCG
CGTTAATATCGCTAG
- How long is the original sequence? Reconstruct it.
- Name a gene variant that you would want to know is in your genome, one that you wouldn't, and one that you consider to be frivolous.
- One newly identified human gene has counterparts (homologs) in bacteria, yeast, roundworms, mustard weed,

fruit flies, mice, and chimpanzees. A second gene has homologs in fruit flies, mice, and chimpanzees only. What does this information suggest about the functions of these two human genes with respect to each other?

6. Headlines about sequencing genomes of such unusual organisms as sea squirts and pufferfish often serve as material for comedians. Why, scientifically, is it important to sequence the genomes of a variety of organisms?

Web Activities

- Invent an "omics". Consult Omics.org for hints.
- Go to the Genetics Alliance website, and describe a disease-specific organization.
- Consult the <http://www.hgvs.org/dblist/glsdb.html> database, and describe three different mutations that cause a particular disorder.
- Go to the ELSI web pages (<http://www.genome.gov/10001618>). Discuss one societal concern arising from genomics, and how it might affect you.
- Go to the website for The Institute For Genomic Research (<http://www.tigr.org/>). List five species whose genomes have been sequenced and the diseases that they cause in humans.

Case Studies and Research Results

12. After the tsunami that devastated countries bordering the Indian Ocean on December 26, 2004, many organisms never before seen washed up on shore. Researchers collected specimens and sequenced DNA to try to classify the fishes. Consider the following 8-base sequence that is similar among the species:

Fish 1	A	A	G	C	T	A	A	T
Fish 2	T	A	G	C	T	A	C	T
Fish 3	G	T	A	C	T	T	G	T
Fish 4	C	C	T	C	G	C	G	T
Position	1	2	3	4	5	6	7	8

- Write the DNA sequences for the two most closely related fishes.
- Which position(s) in the sequence are highly conserved?
- Which position(s) in the sequence are the least conserved?
- Which site is probably not essential, and how do you know this?
- A coelacanth has C T A C T G G T for this section of the genome. Which of the mystery fishes is the coelacanth's closest relative?

Forensics Focus

- This book has discussed many types of genetic information, from the SNPs that represent single bases, to genome-wide patterns of SNPs and copy number variants, to single-gene mutations, to sequenced exomes, to sequenced genomes. Select one of these and discuss how it would provide more detailed information than the traditional approach of identifying individuals on the basis of comparing a few dozen markers.
- On the soap opera "The Young and the Restless," Chloe had sex with half-brothers Billy and Cane within a few days. After the birth, the half-brothers battled over who was the daddy, because the DNA profiles, based on the CODIS markers (see figure 14.10), were identical. Could personal genome sequencing distinguish the feuding fathers sufficiently to establish parenthood? (Cane, it turned out, was an imposter who supplied DNA for his test stolen from Billy's real half-brother.)

Glossary

Pronunciations are provided for technical terms.

KEY ə = eh

— = long vowel sound

˘ = short vowel sound

ˊ = heavy accent

ˋ = light accent

^ = aw

A

acrocentric chromosome (äk' rō sĕn'trĭk krōmā sōm) A chromosome in which the centromere is near one end.

adaptive immunity (ĭ-myōō'nĭ-tē) A slow, specific immune response following exposure to a foreign antigen.

adenine (äd'en-ēn') One of two purine nitrogenous bases in DNA and RNA.

allele (ə-lĕl') An alternate (variant) form of a gene.

alternate splicing Building different proteins by combining exons of a gene in different ways.

amino (ə-mĕ'nō) **acid** A small organic molecule that is a protein building block.

amniocentesis (äm'nĕ-ō-sĕn-tĕ'sĭs) A test that examines fetal chromosomes and biochemicals in amniotic fluid.

anaphase (än-ä-fäz') When the centromeres of replicated chromosomes part in mitosis.

aneuploid (än'yü-plōid') A cell with one or more extra or missing chromosomes.

angiogenesis (än'je-o-jĕn'ō-sĭs) Extension of blood vessels.

antibody (än'tĕ-bōd'ē) A multisubunit protein, produced by B cells, that binds a specific foreign antigen, alerting the immune system or destroying the antigen.

anticodon (äntĕ-kō'dōn) A three-base sequence on one loop of a transfer RNA molecule that is complementary to an mRNA codon, and joins to the appropriate amino acid and its mRNA.

antigen (än'tĕ-jĕn) A molecule that elicits an immune response.

antigen binding sites (än'tĕ-jĕn) Specialized ends of antibody chains.

antigen (än'tĕ-jĕn) -**presenting cell** A cell displaying a foreign antigen.

antiparallel The head-to-tail position of the entwined chains of the DNA double helix.

apoptosis (äpō-tō'sĭs) A form of cell death that is a normal part of growth and development.

assisted reproductive technologies Procedures that replace a gamete or the uterus to help people with fertility problems have children.

autoantibodies (ō'tō-än'tĕ-bōdĕz) Antibodies that attack the body's own cells.

autoimmunity (ō'tō-ĭ-myōō'nĭ-tē) An immune attack against one's own body.

autosomal (ōtə-sōmäl) **dominant** The inheritance pattern of one autosomal allele causing a phenotype, which can affect males and females and does not skip generations.

autosomal (ōtə-sōmäl) **recessive** The inheritance pattern of two autosomal alleles causing a phenotype, which can affect males and females and can skip generations.

autosome (ōtə-sōm) A chromosome that does not have a gene that determines sex.

B

balanced polymorphism (pōl'ĕ-mōr'fizəm) Maintenance of a harmful recessive allele in a population because the heterozygote has a reproductive advantage.

base excision repair Removal of up to five contiguous DNA nucleotides to correct oxidative damage.

B cell A type of lymphocyte that secretes antibody proteins in response to nonself antigens displayed on other immune system cells.

bioethics A field that addresses personal issues that arise in applying medical technology and genetic information.

bioremediation Use of plants or microorganisms to detoxify environmental pollutants.

biotechnology The alteration of cells or biochemicals with a specific application.

blastocyst (bläs'tō-sĭst') A hollow ball of cells descended from a fertilized ovum.

blastomere (blästō-mĕr') A cell of a blastocyst.

C

cancer (kän'sər) A group of disorders resulting from loss of cell cycle control.

cancer stem cells Stem cells that divide and yield cancer cells and abnormal specialized cells.

carbohydrate (kar'bo-hĭ'-drät) An organic compound that consists of carbon, hydrogen, and oxygen in a 1:2:1 ratio. Includes sugars and starches.

carcinogen (kar-sĭn-ō-jĕn) A substance that causes cancer.

case-control study An epidemiological method in which people with a particular condition are compared to individuals as much like them as possible, but without the disease.

cDNA library A collection of DNA molecules reverse transcribed from the mRNAs in a particular cell type.

cell (sel) The fundamental unit of life.

cell (sel) **cycle** A cycle of events describing a cell's preparation for division and division itself.

cellular adhesion A precise series of interactions among the proteins that connect cells.

cellular immune (ĭ-myōōn') **response** T cells release cytokines to stimulate and coordinate an immune response.

centriole (sĕn'-trĕ-ohl) A structure in cells that organizes microtubules into the mitotic spindle.

centromere (sĕn'-trō-mĭr) The largest constriction in a chromosome, located at a specific site in each chromosome type.

centrosome (sĕn'-trō-sōm) A structure built of centrioles and proteins that organizes microtubules into a spindle during cell division.

chaperone protein A protein that binds a polypeptide and guides folding.

chorionic villus (kōrĕ-ōn'ĭk vĭl-us) **sampling** (CVS) A prenatal diagnostic technique that analyzes chromosomes in chorionic villus cells, which, like the fetus, descend from the fertilized ovum.

chromatid (krō' mā-tĭd) A single, very long DNA molecule and its associated proteins, forming a longitudinal half of a replicated chromosome.

chromatin (krō' mā-tĭn) DNA and its associated proteins.

chromatin (krō' mā-tĭn) **remodeling** Adding or removing chemical groups to or from histones, which can alter gene expression.

chromosome (krō' mā-sōm') A highly wound continuous molecule of DNA and the proteins wrapped around it.

chromosome microarray analysis A technique that detects small copy number variants used with amniocentesis or chorionic villus sampling.

cleavage (klĕvĭj) A series of rapid mitotic cell divisions after fertilization.

clines (klĭnz) Allele frequencies that change from one geographical area to another.

cloning vector A piece of DNA used to transfer DNA from a cell of one organism into the cell of another.

coding strand The side of the DNA double helix for a particular gene from which RNA is not transcribed.

codominant A heterozygote in which both alleles are fully expressed.

codon (kō'dōn) A continuous triplet of mRNA that specifies a particular amino acid.

coefficient of relatedness The proportion of genes that two people related in a certain way share.

cohort study An investigation that follows a large group of individuals over time while recording or assessing a health measure.

collectins (ko-lek'tinz) Immune system molecules that detect viruses, bacteria, and yeasts.

comparative genomics (jə-nō'mīks) Identifying conserved DNA sequences among genomes of different species.

comparative genomic hybridization (CGH) A technique using fluorescent labels to detect copy number variants.

complement Plasma proteins that have a variety of immune functions.

complementary base pairs The pairs of DNA bases that hydrogen bond; adenine bonds to thymine and guanine to cytosine.

complementary DNA (cDNA) A DNA molecule that is the complement of an mRNA, copied using reverse transcriptase.

concordance (kən-kōr'dens) A measure indicating the degree to which a trait is inherited; percentage of twin pairs in which both members express a trait.

conditional mutation (myōō-tā'shən) A genotype that is expressed only under certain environmental conditions.

conformation The three-dimensional shape of a molecule.

consanguinity (kōnsān-gwīn'ī-tē) Blood relatives having children together.

copy number variant A DNA sequence present in different numbers of copies in different individuals; repeats.

critical period The time during prenatal development when a structure is sensitive to damage from a mutation or an environmental intervention.

crossing over An event during prophase I when homologs exchange parts.

cytogenetics (sītō-jə-nēt'īks) Matching phenotypes to detectable chromosomal abnormalities.

cytokine (sītō-kīn') A biochemical that a T cell secretes that controls immune function.

cytokinesis (sī-tō-kīn-ē'sis) Division of the cytoplasm and its contents.

cytoplasm (sī'tō-plāzm) Cellular contents other than organelles.

cytosine (sī'tō-sēn) One of the two pyrimidine nitrogenous bases in DNA and RNA.

cytoskeleton (sī-tō-skēl'ī-tn) A framework of protein tubules and rods that supports the cell and gives it a distinctive form.

D

dedifferentiated A cell less specialized than the cell it descends from. A characteristic of a cancer cell.

deletion mutation (myōō-tā'shən) A missing sequence of DNA or part of a chromosome.

deoxyribonucleic acid (dē-ōksē-rībō-nōō-klā'īk) (DNA) The genetic material; the biochemical that forms genes.

deoxyribose (dē-ōksē-rī'bōs) 5-carbon sugar in a DNA nucleotide.

differentiation Cell specialization, reflecting differential gene expression.

dihybrid cross Breeding individuals heterozygous for two traits.

diploid (dīp'loid) A cell containing two sets of chromosomes.

dizygotic (dīzī-gōt'īk) (**DZ**) **twins** Twins that originate as two fertilized ova; fraternal twins.

DNA See **deoxyribonucleic acid**.

DNA damage response DNA repair.

DNA microarray See **microarray**.

DNA polymerase (pə-līm'ər-ās) (DNAP) An enzyme that adds new bases to replicating DNA and corrects mismatched base pairs.

DNA probe A labeled short sequence of DNA that binds its complement in a biological sample.

DNA profiling A biotechnology that detects differences in the number of copies of certain DNA repeats among individuals. Used to rule out or establish identity.

DNA replication Construction of a new DNA double helix using the information in parental strands as a template.

dominant A gene variant expressed when present in one copy.

duplication An extra copy of a DNA sequence, usually caused by misaligned pairing in meiosis.

E

ectoderm (ēktō-dūrm) The outermost primary germ layer.

embryo (ēm'brē-ō') In humans, prenatal development until the end of the eighth week, when all basic structures are present.

embryonic (ēmbrē-ōn'īk) **stem (ES) cell** A cell, derived in laboratory culture from inner cell mass cells of very early embryos, that can self-renew and differentiate as any cell type.

empiric risk Probability that a trait will recur based upon its incidence in a population.

endoderm (ēn'dō-dūrm) The innermost primary germ layer of the primordial embryo.

endoplasmic reticulum (ēndō-plāzmīk rə-tīk'u-lōm) (ER) An organelle consisting of a labyrinth of membranous tubules on which proteins, lipids, and sugars are synthesized.

enzyme (ēnzīm) A type of protein that speeds the rate of a specific biochemical reaction.

epigenetic (ēpē-jə-nēt'īk) A layer of information placed on a gene that is a modification other than a change in DNA sequence, such as methylation.

epistasis (ēpē-stā-sis) A gene masking the expression of another.

epitope (ep'ī-tōp) Part of an antigen that an antibody binds.

equational division The second meiotic division, producing four cells from two.

euchromatin (yōō-krō'mā-tīn') Parts of chromosomes that do not stain and that contain active genes.

eugenics (yōō-jēn'īks) The control of individual reproductive choices to achieve a societal goal.

eukaryotic cell (yōō-kar'ē-ōt'īk sel) A complex cell containing organelles, including a nucleus.

euploid (yōō'plōid) A somatic cell with the normal number of chromosomes for the species.

excision repair Enzyme-catalyzed removal of pyrimidine dimers in DNA.

exome (x-ōm) The part of the genome that encodes protein.

exon (x-on) Part of a gene that encodes amino acids.

expressivity Degree of severity of a phenotype.

F

fetus (fē'təs) The prenatal human after the eighth week of development, when structures grow and specialize.

founder effect A type of genetic drift in which a few individuals found a new settlement, perpetuating a subset of alleles from the original population.

frameshift mutation (myōō-tā'shən) A mutation that alters a gene's reading frame.

fusion protein A protein that forms from translation of transcripts from two genes.

G

G₀ An offshoot of the cell cycle in which the cell remains specialized but does not replicate its DNA or divide.

G₁ The stage of the cell cycle following mitosis in which the cell resumes synthesis of proteins, lipids, and carbohydrates.

G₂ The stage of the cell cycle following S phase but before mitosis, when certain proteins are synthesized.

gamete (gām'ēt) A sex cell.

gamete intrafallopian transfer (GIFT) An infertility treatment in which sperm and oocytes are placed in a woman's uterine tube.

gastrula (gāstrə-lə) A three-layered embryo.

gene (jēn) A sequence of DNA that instructs a cell to produce a particular protein.

gene expression Transcription of a gene's DNA into RNA.

gene expression profiling Use of DNA microarrays to detect the types and amounts of cDNAs reverse transcribed from the mRNAs in a particular cell source.

gene pool All the genes in a population.

gene therapy Replacing a malfunctioning gene to correct an abnormality.

genetic (jə-nēt'īk) **code** The correspondence between specific mRNA triplets and the amino acids they specify.

genetic counselor A medical specialist who calculates risk of recurrence of inherited disorders in families, applying the laws of inheritance to pedigrees.

genetic determinism Attributing a trait entirely to a gene or genes.

genetic drift Changes in allele frequencies in small groups reproductively separated from a larger population.

genetic heterogeneity A phenotype that can be caused by variants of any of several genes.

genetic load The collection of deleterious recessive alleles in a population.

genetic marker DNA sequences near genes of interest that are co-inherited unless separated

by a crossover. SNPs and copy number variants are used as markers.

genetics The study of inherited variation.

genome (jē'nōm) The complete set of genetic instructions in the cells of a particular type of organism.

genome-wide association study A case-control study in which many thousands of variants (single nucleotide polymorphisms or copy number variants) that form haplotypes are compared between people with a condition and unaffected individuals.

genomic (jē nōm'ik) **imprinting** Differing of the phenotype depending upon which parent transmits a particular allele.

genomic (jē nōm'ik) **library** A collection of DNA pieces representing the genome of an individual, including introns.

genomics (jē nōm'iks) The study of the functions and interactions of many genes or other DNA sequences, or comparing genomes.

genotype (jē n'ə- tīp) The allele combinations in an individual that cause particular traits or disorders.

genotypic (jēn'ə- tīp'ik) **ratio** The ratio of genotype classes expected in the progeny of a particular cross.

germline gene therapy Genetic alterations of gametes or fertilized ova, which perpetuate the change throughout the organism and transmit it to future generations.

germline mutation A mutation in every cell in an individual.

Golgi (gōl'jē) **apparatus** An organelle, consisting of flattened, membranous sacs, that packages secretion components.

gonads (gō'nadz) Paired structures in the reproductive system where sperm or oocytes are manufactured.

growth factor A protein that stimulates mitosis.

guanine (gwa'nēn) One of the two purine nitrogenous bases in DNA and RNA.

H

haploid (hăp'loid) A cell with one set of chromosomes.

haplogroup (hăp' lō-grōp) In DNA ancestry testing, a specific set of markers on the Y or mitochondrial chromosome.

haplotype (hăp' lō tīp) A series of known DNA sequences or single nucleotide polymorphisms linked on a chromosome.

Hardy-Weinberg equilibrium An idealized state in which allele frequencies in a population do not change from generation to generation.

heavy chains The two longer polypeptide chains of an antibody subunit.

hemizygous (hēm' ē-zī' gəs) The sex that has half as many X-linked genes as the other; a human male.

heritability An estimate of the proportion of phenotypic variation in a group due to genes.

heterochromatin (hēt'ə-rō-krō'mā tīn) Dark-staining chromosome parts that have few protein-encoding genes.

heterogametic (hēt'ə-rō-gə-mē'tīk) **sex** The sex with two different sex chromosomes; a human male.

heteroplasmy (hēt'ə-rō-plāz-mē) Mitochondria in the same cell having different alleles of a particular gene.

heterozygous (hēt'ə-rō-zī' gəs) Having two different alleles of a gene.

histone (hīs'tōn) A type of protein around which DNA entwines.

hominins (hōm'ə-nīnz) Animals ancestral to humans only.

hominoids (hōm'ə-nōīdz) Animals ancestral to apes and humans only.

homogametic (hō'mō-gə-mē'tīk) **sex** The sex with identical types of sex chromosomes; the human female.

homologous (hō-mōl'ə-gəs) **pairs** Chromosomes with the same gene sequence.

homozygosity (hōmō-zī-gəs'ī-tē) **mapping** An approach to gene discovery that correlates stretches of homozygous DNA base sequence in the genomes of related individuals to certain traits or disorders.

homozygous (hōmō-zī' gəs) Having two identical alleles of a gene.

hormone (hor' mōn) A biochemical produced in a gland and carried in the blood to a target organ, where it exerts an effect.

human leukocyte antigen (lōōkə-sīt' ən'ti-jən) (HLA) **complex** Genes closely linked on the short arm of chromosome 6 that encode cell surface proteins important in immune system function.

humoral (yōō' mər-əl) **immune response** Process in which B cells secrete antibodies into the bloodstream.

I

idiotype (id'ē-o-tīp) Part of an antibody molecule that binds an antigen.

incidence The number of new cases of a disease during a certain time in a particular population.

incomplete dominance A heterozygote intermediate in phenotype between either homozygote.

independent assortment The random arrangement of homologous chromosome pairs, in terms of maternal or paternal origin, down the center of a cell in metaphase I. Inheritance of a gene on one chromosome does not influence inheritance of a gene on a different chromosome. (Mendel's second law)

induced pluripotent stem (iPS) cells Somatic cells that are genetically reprogrammed to differentiate differently.

infertility The inability to conceive a child after a year of unprotected intercourse.

inflammation Part of the innate immune response that causes an infected or injured area to swell with fluid, turn red, and attract phagocytes.

innate immunity (ī-myōō'nī-tē) Components of immune response that are present at birth and do not require exposure to an environmental stimulus.

inner cell mass A clump of cells on the inside of the blastocyst that will continue developing into an embryo.

insertional translocation A rare type of translocation in which a part of one chromosome is part of a nonhomologous chromosome.

insertion mutation (myōō-tā'-shən) A mutation that adds DNA bases.

interferon (in'tər-fēr'on) A type of cytokine.

interleukin (in'tər-loo'kin) A type of cytokine.

intermediate filament A type of cytoskeletal component made of different proteins in different cell types.

interphase (in'tər-fāz') Stage when a cell is not dividing.

intracytoplasmic (in'trə-sītō-plāzmīk) **sperm injection** (ICSI) An infertility treatment that injects a sperm cell nucleus into an oocyte, to overcome lack of sperm motility.

intrauterine (in'trə-yōō'tər-in) **insemination** An infertility treatment that places donor sperm in the cervix or uterus.

intron (in trōn) Part of a gene that is transcribed but is excised from the mRNA before translation into protein.

in vitro (in vē'trō) **fertilization** (IVF) Placing oocytes and sperm in a laboratory dish with appropriate biochemicals so that fertilization occurs, then, after a few cell divisions, transferring the embryos to a woman's uterus.

in vivo gene therapy Direct genetic manipulation of cells in the body.

K

karyotype (kā'rē-ō-tīp) A size-order chromosome chart.

L

law of independent assortment See **independent assortment**.

law of segregation See **segregation**.

lethal allele (ə-lēl') An allele that causes death before reproductive maturity or halts prenatal development.

ligand (lī'gənd) A molecule that binds to a receptor.

ligase (lī'gās) An enzyme that catalyzes the formation of covalent bonds in the sugar-phosphate backbone of a nucleic acid.

light chains The two shorter polypeptide chains of an antibody subunit.

linkage Genes on the same chromosome.

linkage disequilibrium Extremely tight linkage between DNA sequences.

linkage maps Maps that show gene order on chromosomes, determined from crossover frequencies between pairs of genes.

lipid (lipīd) A type of organic molecule that has more carbon and hydrogen atoms than oxygen atoms. Includes fats and oils.

lysosome (lī'sō-sōm) A saclike organelle containing enzymes that degrade debris.

M

macroevolution (māk'rō-ēv'ə-lōōshən) Genetic change sufficient to form a new species.

major histocompatibility complex (hīstō-kəm-pātə-bilī-tē) (MHC) A gene cluster, on chromosome 6 in humans, that includes many genes that encode components of the immune system.

manifesting heterozygote (hēt'ə-rō-zīgōt) A female carrier of an X-linked recessive gene who expresses the phenotype because the normal allele is inactivated in some tissues.

meiosis (mī-ō'sīs) Cell division that halves the number of chromosomes to form haploid gametes.

memory cells B or T cell descendants that carry out a secondary immune response.

mesoderm (mēz-ō-dūrm) The middle primary germ layer.

messenger RNA (mRNA) A molecule of RNA complementary in sequence to the template strand of a gene that specifies a protein product.

metacentric chromosome (mētə-sēn'trīk krōmōsōm) A chromosome with the centromere approximately in the center.

metaphase (mētə-fāz) The stage of mitosis when chromosomes align along the center of the cell.

metastasis (mētə-stā'sīs) Spread of cancer from its site of origin to other parts of the body.

microarray A set of target genes embedded in a glass chip, to which labeled cDNAs from a sample bind and fluoresce. Microarrays show patterns of gene expression.

microevolution Change of allele frequency in a population.

microfilament A solid rod of actin protein that forms part of the cytoskeleton.

microRNA A 21- or 22-base-long RNA that binds to certain mRNAs, blocking their translation into protein.

microtubule (mīkrō-tōōbyōōl) A hollow structure built of tubulin protein that forms part of the cytoskeleton.

mismatch repair Proofreading of DNA for misalignment of short, repeated segments.

missense (mīs'sēns) A single base change mutation that alters an amino acid.

mitochondrion (mītō-kōn'drē-ən) An organelle consisting of a double membrane that houses enzymes that catalyze reactions that extract energy from nutrients.

mitosis (mī-tō'sīs) Division of somatic (nonsex) cells.

mode of inheritance The pattern in which a gene variant passes from generation to generation. It may be dominant or recessive, autosomal or sex-linked.

modifier gene A gene that affects expression of another gene.

molecular evolution Changes in protein and DNA sequences over time used to estimate how recently species diverged from a shared ancestor.

monohybrid (mōn'ō-hībrīd) **cross** A cross of two individuals who are heterozygous for a single trait.

monosomy (mōn'ō-sō'mē) A human cell with 45 (one missing) chromosomes.

monozygotic (mōnō-zī-gōt'īk) (MZ) **twins** Twins that originate as a single fertilized ovum; identical twins.

morula (mōr' yə-lə) The very early prenatal stage that resembles a mulberry.

multifactorial trait A trait or illness determined by several genes and the environment.

mutagen (myōō'tə-jən) A substance that changes, adds, or deletes a DNA base.

mutant (myōō'tnt) An allele that differs from the normal or most common allele in a population that alters the phenotype.

mutation (myōō-tā'shən) A change in a protein-encoding gene that affects the phenotype and occurs in less than one percent of a population.

N

natural selection Differential survival and reproduction of individuals with particular phenotypes in particular environments, which may alter allele frequencies in subsequent generations.

neural (nōō'r'əl) **tube** A structure in the embryo that develops into the brain and spinal cord.

nitrogenous (nī-trōj'ə-nəs) **base** A nitrogen-containing base that is part of a nucleotide.

nondisjunction (nōndīs-jūngk'shən) The unequal partitioning of chromosomes into gametes during meiosis.

nonsense mutation (myōō'tā'shən) A point mutation that changes an amino-acid-coding codon into a stop codon, prematurely terminating synthesis of the encoded protein.

non synonymous codon (kō'don) A codon that encodes a different amino acid from another codon.

nucleic (nōō-klē'īk) **acid** DNA or RNA.

nucleolus (nōō-klē'ō-ləs) A structure in the nucleus where ribosomes are assembled from ribosomal RNA and protein.

nucleosome (nōō-klē'ō-sōm) A unit of chromatin structure.

nucleotide (nōō-klē'ō-tīd) The building block of a nucleic acid, consisting of a phosphate group, a nitrogenous base, and a 5-carbon sugar.

nucleotide (nōō-klē'ō-tīd) **excision repair** Replacement of up to 30 nucleotides to correct DNA damage of several types.

nucleus (nōō-klē-əs) A large, membrane-bounded region of a eukaryotic cell that houses DNA.

oncogene (ōn'kə-jēn) A gene that normally controls the cell cycle, but causes cancer when overexpressed.

oocyte (ō'və-sīt) The female gamete (sex cell).

oogenesis (ōə-jēn'ī-sīs) Oocyte development.

organelle (ōr'gə-nēl') A specialized structure in a eukaryotic cell that carries out a specific function.

ovaries (ō'və-rēz) The female gonads.

P

paracentric (para sēn'trīk) **inversion** An inverted chromosome that does not include the centromere.

pedigree A chart of symbols connected by lines that depict the genetic relationships and transmission of inherited traits in related individuals.

penetrance Percentage of individuals with a genotype who have an associated phenotype.

pericentric (pēr-ē sēn-trīk) **inversion** An inverted chromosome that includes the centromere.

peroxisome (pə-rōk'sī-sōm) An organelle consisting of a double membrane that houses enzymes with various functions.

pharmacogenetics (farm a kōjə-nēt-īks) Testing for single gene variants that affect drug metabolism.

pharmacogenomics (farm a kōjə-nōm-īks) Testing for variants of many genes or gene expression profiles that affect drug metabolism.

phenocopy (fē' nō-kōp'ē) An environmentally caused trait that occurs in a familial pattern, mimicking inheritance.

phenotype (fē' nō-tīp) The expression of a gene in traits or symptoms.

plasma cell A cell descended from a B cell that produces abundant antibodies of a single type.

plasma membrane (plāz'mə mēm'brān) The selective barrier around a cell, consisting of proteins, glycolipids, glycoproteins, and lipid rafts on or in a phospholipid bilayer.

plasmid (plāz' mīd) A small circle of double-stranded DNA found in some bacteria. Used as a vector in recombinant DNA technology.

pleiotropic (plēə-trōpīk) A single-gene disorder with several symptoms. Different symptom subsets may occur in different individuals.

point mutation (myōō-tā' shən) A single base change in DNA.

polar body A product of female meiosis that contains little cytoplasm and does not continue to develop into an oocyte.

polar body biopsy (bī'ōp sē) A genetic test performed on a polar body to infer the genotype of the attached oocyte.

polygenic (pōlē-jēn' īk) **traits** Traits determined by more than one gene.

polymerase (pōl'ə-mə-rās) **chain reaction** (PCR) A nucleic acid amplification technique in which a DNA sequence is replicated in a test tube to rapidly produce many copies.

polymorphism (pōlē-mōr' fīz əm) A DNA base or sequence at a certain chromosomal locus that varies in at least 1 percent of individuals in a population.

polyploid (pŏl'ē-ploid) A cell with one or more extra sets of chromosomes.

population A group of interbreeding individuals.

population bottleneck Decrease in allele diversity resulting from an event that kills many members of a population, followed by restoration of population numbers.

population genetics (jə-nĕt'ĭk) The study of allele frequencies in different groups of individuals.

population study Comparison of disease incidence in different groups of people.

preimplantation genetic (4th) diagnosis (PGD) Removing a cell from an 8-celled embryo and testing it for a mutation to deduce the genotype of the embryo.

prevalence The number of cases of a disease in a population at a particular time.

primary germ layers The three layers of an embryo.

primary immune response Immune system's response to initial encounter with a nonself antigen.

primary (1st) structure The amino acid sequence of a protein.

progenitor cell A cell whose descendants can follow any of several developmental pathways.

prokaryotic cell (prŏ-kārĕ-ŏt'ĭk sĕl) A cell that does not have a nucleus or other organelles. One of the three domains of life. Bacteria.

promoter A control sequence near the start of a gene.

pronuclei (prŏ-nŏŏ'klĕĭ) DNA packets in the fertilized ovum.

prophase (prŏ'fāz) The first stage of mitosis or meiosis, when chromatin condenses.

prospective study A study that follows two or more groups.

proteasome (prŏ-tĕ-ə-sŏm) A multiprotein structure in a cell shaped like a barrel through which misfolded proteins pass and are refolded or dismantled.

protein A type of macromolecule that is the direct product of genetic information; a chain of amino acids.

proteome (prŏ'tĕ-ŏm) The set of proteins a cell produces.

proteomics (prŏtĕ-ŏ' mĭk) Study of the proteins produced in a particular cell type under particular conditions.

proto-oncogene (prŏtŏ-ŏn'kŏ-jĕn) A gene that normally controls the cell cycle, but when overexpressed causes cancer.

pseudogene (sŏŏ' dŏ-jĕn) A gene that does not encode protein, but whose sequence very closely resembles that of a coding gene.

Punnett square A diagram used to follow parental gene contributions to offspring.

purine (pyŏŏ'rĕn) A DNA base with a two-ring structure; adenine and guanine are purines.

pyrimidine (pĭ-rĭm'ĭ-dĕn) A DNA base with a single-ring structure; cytosine, thymine, and uracil are pyrimidines.

Q

quantitative trait loci Genes that determine polygenic traits.

quaternary (4th) structure A protein that has more than one polypeptide subunit.

R

reading frame The grouping of DNA base triplets encoding an amino acid sequence.

receptor A structure on a cell that binds a specific molecule.

recessive An allele whose expression is masked by another allele.

reciprocal translocation A chromosome aberration in which two nonhomologous chromosomes exchange parts, conserving genetic balance but rearranging genes.

recombinant (rĕ-kŏm'bŏ-nŏnt) A series of alleles on a chromosome that differs from the series of either parent.

recombinant (rĕ-kŏm'bŏ-nŏnt) DNA technology Transferring genes between species.

reduction division The first meiotic division, which halves the chromosome number.

replication fork Locally opened portion of a replicating DNA double helix.

ribonucleic acid (RNA) (rĭ bŏ-nŏŏ-klĕ'ĭk) A nucleic acid whose bases are A, C, U, and G.

ribose (rĭ'bŏs) A 5-carbon sugar in RNA.

ribosomal (rĭ'bŏs-ŏ'mŏl) **RNA (rRNA)** RNA that, with proteins, comprises ribosomes.

ribosome (rĭ'bŏsŏm) An organelle consisting of RNA and protein that is a scaffold for protein synthesis.

risk factor A characteristic or experience associated with increased likelihood of developing a particular medical condition.

RNA interference Introduction of a small interfering RNA molecule that binds to and prevents translation of a specific mRNA.

RNA polymerase (RNAP) (pŏl'ə-mə-rās) An enzyme that adds RNA nucleotides to a growing RNA chain.

Robertsonian (Răb-ŏrt - sŏ-nĕ-ŏn)
translocation A chromosome aberration in which two short arms of nonhomologous chromosomes break and the long arms fuse, forming one unusual, large chromosome.

run of homozygosity Regions of the genome in which contiguous SNPs (single nucleotide polymorphisms) are homozygous, indicating a shared ancestor with another person with the same pattern.

S

S phase The stage of interphase when DNA replicates.

secondary immune response Immune system activation in response to a second or subsequent encounter with a pathogen.

secondary (2nd) structure Folds in a polypeptide caused by attractions between amino acids close together in the primary structure.

segregation The distribution of alleles of a gene into separate gametes during meiosis. (Mendel's first law).

self-renewal Defining property of a stem cell; the ability to yield a daughter cell like itself.

semiconservative replication DNA synthesis along each half of the double helix.

sex chromosome (krŏ'mŏ-sŏm) A chromosome containing genes that specify sex.

sex-influenced trait Phenotype caused when an allele is recessive in one sex but dominant in the other.

sex-limited trait A trait that affects a structure or function present in only one sex.

sex ratio Number of males divided by number of females multiplied by 1,000 for people of a certain age in a population.

short tandem repeats (STRs) Repeats of 2 to 10 DNA bases that are compared in DNA profiling.

signal transduction A series of biochemical reactions and interactions that pass information from outside a cell to inside, triggering a response.

single nucleotide polymorphism (nŏŏklĕŏ-tĭd pŏlĕ-mŏr' fĭz'ŏm) (SNP) Single base sites that differ among individuals. A SNP is present in at least 1 percent of a population.

somatic cell (sŏ-măt'ĭk sĕl) A nonsex cell, with 23 pairs of chromosomes in humans.

somatic (sŏ-măt'ĭk) gene therapy Genetic alteration of a specific cell type, not transmitted to future generations.

somatic mutation (sŏ-măt'ĭk myŏŏ-tă'shŏn) A genetic change in a nonsex cell.

spermatogenesis (spŏr-măt'ə-jĕn'ĭ-sis) Sperm cell differentiation.

spermatogonium (spŏr'mah-to-gŏ ne-um) An undifferentiated cell in a seminiferous tubule that can give rise to a sperm cell in meiosis.

spermatozoon (spŏr-măt'ə-zŏ'ŏn) (sperm) A mature male reproductive cell (meiotic product).

spindle A structure composed of microtubules that pulls sets of chromosomes apart in a dividing cell.

spontaneous mutation (myŏŏ-tă'sheŏn) A genetic change that results from mispairing when the replication machinery encounters a base in its rare tautomeric form.

SRY gene The sex-determining region of the Y. If the SRY gene is activated, the gonad develops into a testis; if not, an ovary forms under direction of other genes.

stem cells Cells that give rise to other stem cells, as well as to cells that differentiate.

submetacentric chromosome (sŭb mĕt-ə-sĕn'trĭk krŏ'mŏ-sŏm) A chromosome in which the centromere establishes a long arm and a short arm.

sugar-phosphate backbone The "rails" of a DNA double helix, consisting of alternating deoxyribose and phosphate groups, oriented opposite one another.

synonymous codons (kŏd ŏnz) DNA triplets that specify the same amino acid.

synteny (sĭn'tŏ-nĕ) Correspondence of genes on the same chromosome in several species.

T

- tandem duplication** A duplicated DNA sequence next to the original sequence.
- T cell** A type of lymphocyte that produces cytokines and coordinates the immune response.
- telomerase** (tə-lōm'ə-rās) An enzyme, including a sequence of RNA, that adds DNA to chromosome tips.
- telomere** (tēl'ə-mîr) A chromosome tip.
- telophase** (tēl'ə-fāz) The stage of mitosis or meiosis when daughter cells separate.
- template strand** The DNA strand carrying the information to be transcribed.
- teratogen** (tə-rāt'ə-jən) A substance that causes a birth defect.
- tertiary (3°) structure** Folds in a polypeptide caused by interactions between amino acids and water. This draws together amino acids that are far apart in the primary structure.
- testes** (tes'tēz) The male gonads.
- thymine** (thī'mēn) One of the two pyrimidine bases in DNA.
- transcription** Manufacturing RNA from DNA.
- transcription factor** A protein that activates the transcription of certain genes.
- transfer RNA (tRNA)** A type of RNA that connects mRNA to amino acids during protein synthesis.
- transgenic organism** (trāns-jěn'ĭk) An individual with a genetic modification in every cell.
- transition** A point mutation altering a purine to a purine or a pyrimidine to a pyrimidine.

- translation** Assembly of an amino acid chain according to the sequence of base triplets in a molecule of mRNA.
- translocation** Exchange between nonhomologous chromosomes.
- translocation carrier** An individual with exchanged chromosomes but no signs or symptoms. The person has the usual amount of genetic material, but it is rearranged.
- transposon** (trāns-pōzōn) A gene or DNA segment that moves to another chromosome.
- transversion** A point mutation altering a purine to a pyrimidine or vice versa.
- trisomy** (trī sō'mē) A human cell with 47 chromosomes (one extra).
- tumor suppressor gene** (tōmər səprēs'ər jēn) A recessive gene whose normal function is to limit the number of divisions a cell undergoes.

U

- uniparental disomy** (yū-nī-pə'rent-əl dī sō mē) Inheriting two copies of the same gene from one parent.
- uracil** (yōōr'ə-sīl) One of the four types of bases in RNA; a pyrimidine.

V

- vaccine** (vak-sē'n) An inactive or partial form of a pathogen that stimulates antibody production.
- variable number of tandem repeats (VNTRs)** Repeats of 10 to 80 DNA bases that are compared in DNA profiles.

- vesicles** (ves-ə-kulz) Bubble-like membrane-bounded organelles that participate in secretion.
- virus** (vī rəs) An infectious particle built of nucleic acid in a protein coat.

W

- wild type** The most common phenotype in a population for a particular gene.

X

- X inactivation** The inactivation of one X chromosome in each cell of a female mammal, occurring early in embryonic development.
- X-linked** Genes on an X chromosome.
- X-Y homologs** (hōm'ə-lōgz) Y-linked genes that are similar to genes on the X chromosome.

Y

- Y-linked** Genes on a Y chromosome.

Z

- zygote** (zī'gōt) A prenatal human from the fertilized ovum stage until formation of the primordial embryo, at about two weeks.
- zygote intrafallopian transfer** (zī' gōt in'trə-fə-lō' pē-ən) (ZIFT) An assisted reproductive technology in which an ovum fertilized *in vitro* is placed in a woman's uterine tube.

Credits

Line Art and Text

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